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METHOD FOR MODIFYING PLANT MORPHOLOGY,

BIOCHEMISTRY AND PHYSIOLOGY

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TRANSMITTAL OF PRIORITY DOCUMENT

Sir:

Enclosed is a certified copy of European Patent Appln. No.

01870053.4 filed on March 16, 2001, from which priority is claimed under 35 U.S.C. §119.

Respectfully submitted,

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Attestation

Die angehefteten Unterlagen stimmen mit der ursprünglich eingereichten Fassung der auf dem nächsten Blatt bezeichneten europäischen Patentanmeldung überein.

The attached documents are exact copies of the European patent application described on the following page, as originally filed.

Les documents fixés à cette attestation sont conformes à la version initialement déposée de la demande de brevet européen spécificée à la page suivante.

Patentanmeldung Nr.

Patent application No.

Demande de brevet n°

01870053.4 / EP01870053

The organization code and number of your priority application, to be used for filing abroad under the Paris Convention, is EP01870053

CERTIFIED COPY OF PRIORITY DOCUMENT

Der Präsident des Europäischen Patentamts; Im Auftrag

For the President of the European Patent Office Le President de l'Office européen des brevets

R.C. van Dijk

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Bezeichnung der Erfindung/Title of the invention/Titre de l'invention: (Falls die Bezeichnung der Erfindung nicht angegeben ist, siehe Beschreibung. If no title is shown please refer to the description. Si aucun titre n'est indiqué se referer à la description.)

Method for modifying plant morphology, biochemitry and physiology

In anspruch genommene Priorität(en) / Priority(ies) claimed / Priorité(s) revendiquée(s) Staat/Tag/Aktenzeichen / State/Date/File no. / Pays/Date/Numéro de dépôt:

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METHOD FOR MODIFYING PLANT MORPHOLOGY, BIOCHEMISTRY AND PHYSIOLOGY

FIELD OF THE INVENTION

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The present invention generally relates to a method for modifying plant morphological, biochemical and physiological properties or characteristics, such as one or more developmental processes and/or environmental adaptive processes, including but not limited to the modification of initiation or stimulation or enhancement of root growth, and/or adventitious root formation, and/or lateral root formation, and/or root geotropism, and/or shoot growth, and/or apical dominance, and/or branching, and/or timing of senescence, and/or timing of flowering, and/or flower formation, and/or seed development, and/or seed yield, said method comprising expressing a cytokinin degradation control protein, in particular cytokinin oxidase, in the plant, operably under the control of a regulatable promoter sequence such as a cell-specific promoter, tissuespecific promoter, or organ-specific promoter sequence. Preferably, the characteristics modified by the present invention are cytokinin-mediated and/or auxin-mediated characteristics. The present invention extends to genetic constructs which are useful for performing the inventive method and to transgenic plants produced therewith having altered morphological and/or biochemical and/or physiological properties compared to their otherwise isogenic counterparts.

BACKGROUND OF THE INVENTION

Roots are an important organ of higher plants. Their main functions are anchoring of the plant in the soil and uptake of water and nutrients (N-nutrition, minerals, etc.). Thus, root growth has a direct or indirect influence on growth and yield of aerial organs, particularly under conditions of nutrient limitation. Roots are also relevant for the production of secondary plant products, such as defense compounds and plant hormones.

Roots are also storage organs in a number of important staple crops. Sugar beet is the most important plant for sugar production in Europe (260 Mill t/year; 38 % of world production). Manioc (cassava), yams and sweet potato (batate) are important starch producers (app. 150 Mill t/year each). Their content in starch can be twice as high as that of potato. Roots are also the relevant organ for consumption in a number of vegetables (e.g. carrots, radish), herbs (e.g. ginger, kukuma) and medicinal plants (e.g. ginseng). In addition, some of the secondary plant products found in roots are of

economic importance for the chemical and pharmaceutical industry. An example is yams, which contain basic molecules for the synthesis of steroid hormones. Another example is shikonin, which is produced by the roots of *Lithospermum erythrorhizon* in hairy root cultures. Shikonin is used for its anti-inflammatory, anti-tumour and wound-healing properties.

Moreover, improved root growth of crop plants will also enhance competitiveness with weedy plants and will improve growth in arid areas, by increasing water accessibility and uptake.

Improved root growth is also relevant for ecological purposes, such as bioremediation and prevention/arrest of soil erosion.

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Root architecture is an area that has remained largely unexplored through classical breeding, because of difficulties with assessing this trait in the field. Thus, biotechnology could have significant impact on the improvement of this trait, because it does not rely on large-scale screenings in the field. Rather, biotechnological approaches require a basic understanding of the molecular components that determine a specific characteristic of the plant. Today, this knowledge is only fragmentary, and as a consequence, biotechnology was so far unable to realize a break-through in this area.

A well-established regulator of root growth is auxin. Application of indole-3-acetic acid (IAA) to growing plants stimulates lateral root development and lateral root elongation (Torrey, Am J Bot 37: 257-264, 1950; Blakely *et al.*, Bot Gaz 143: 341-352, 1982; Muday and Haworth, Plant Physiol Biochem 32: 193-203, 1994). Roots exposed to a range of concentrations of IAA initiated increasing numbers of lateral roots (Kerk *et al.*, Plant Physiol, 122: 925-932, 2000). Furthermore, when roots that had produced laterals in response to a particular concentration of exogenous auxin were subsequently exposed to a higher concentration of IAA, numerous supernumerary lateral roots spaced between existing ones were formed (Kerk *et al.*, Plant Physiol, 122: 925-932, 2000). Conversely, growth of roots on agar containing auxin-transport inhibitors, including NPA, decreases the number of lateral roots (Muday and Haworth, Plant Physiol Biochem 32: 193-203, 1994).

Arabidopsis mutants containing increased levels of endogenous IAA have been isolated (Boerjan et al., Plant Cell 7: 1405-141, 1995; Celenza et al., Gene Dev 9: 2131-2142, 1995; King et al., Plant Cell 7: 2023-2037, 1995; Lehman et al., Cell 85: 183-194, 1996). They are now known to be alleles of a single locus located on chromosome 2. These

mutant seedlings have excess adventitious and lateral roots, which is in accordance with the above-described effects of external auxin application.

The stimulatory effect of auxins on adventitious and lateral root formation suggests that overproduction of auxins in transgenic plants is a valid strategy for increasing root growth. Yet, it is also questionable whether this would yield a commercial product with improved characteristics. Apart from its stimulatory effect on adventitious and lateral root formation, auxin overproduction triggers other effects, such as reduction in leaf number, abnormal leaf morphology (narrow, curled leaves), aborted inflorescences, increased apical dominance, adventitios root formation on the stem, most of which are undesirable from an agronomic perspective (Klee et al., Genes Devel 1: 86-96, 1987; Kares et al., Plant Mol Biol 15: 225-236, 1990). Therefore, the major problem with approaches that rely on increased auxin synthesis is a problem of containment, namely to confine the effects of auxin to the root. This problem of containment is not likely overcome by using tissue-specific promoters: auxins are transported in the plant and their action is consequently not confined to the site of synthesis. Another issue is whether auxins will always enhance the total root biomass. For agar-grown plants, it has been noticed that increasing concentrations progressively stimulated lateral root formation but concurrently inhibited the outgrowth of these roots (Kerk et al., Plant Physiol, 122: 925-932, 2000).

WO 99/06571 discloses a Zea mays cytokinin oxidase gene and the protein encoding the same as well as transgenic plants comprising the same.

The above-mentioned problems related to containment of auxin effects and to maintenance of root outgrowth are solved by the embodiments in the patent claim.

SUMMARY OF THE INVENTION

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The present invention relates to a genetic construct comprising a gene encoding a protein with cytokinin oxidase activity from *Arabidopsis thaliana*. This gene is expressed under control of a regulated promoter. This promoter may be regulated by endogenous tissue-specific or environment-specific factors or, alternatively, it may be induced by application of specific chemicals.

The present invention also relates to a cell or plant containing the genetic construct.

The present invention also relates to a method to modify root architecture and biomass by expression of a cytokinin oxidase gene under control of a promoter that is specific to the root or to certain tissues or cell types of the root.

DETAILED DESCRIPTION OF THE INVENTION

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To by-pass above-mentioned problems associated with increasing auxin biosynthesis, it was decided to follow an alternative approach. We reasoned that down-regulation of auxin antagonists could evoke similar or even superior effects on root growth as compared to increasing auxin levels. Hormone actions and interactions are extremely complex, but we hypothesized that cytokinins could function as auxin antagonists with respect to root growth. Hormone studies on plant tissue cultures have shown that the ratio of auxin versus cytokinin is more important for organogenesis than the absolute levels of each of these hormones, which indeed indicates that these hormones function as antagonists — at least in certain biological processes. Furthermore, lateral root formation is inhibited by exogenous application of cytokinins. Interestingly, also root elongation is negatively affected by cytokinin treatment, which suggests that cytokinins control both root branching and root outgrowth.

Together, current literature data indicate that increasing cytokinin levels negatively affects root growth, but the mechanisms underlying this process are not understood. The sites of cytokinin synthesis in the plant are root tips and young tissues of the shoot. Endogenous concentrations of cytokinins are in the nM range. However, as their quantification is difficult, rather large tissue amounts need to be extracted and actual local concentrations are not known. Also the subcellular compartmentation of cytokinins is not known. It is generally thought that the free base and ribosides are localized in the cytoplasm and nucleus, while glucosides are localized in the vacuole. There exist also different cytokinins with slightly different chemical structure. As a consequence, it is not known whether the effects of exogenous cytokinins should be ascribed to a raise in total cytokinin concentration or rather to the competing out of other forms of plant-borne cytokinins (which differ either in structure, cellular or subcellular location) for receptors, translocators, transporters, modifying enzymes ...

In order to test the hypothesis that cytokinin levels in the root indeed exceed the level optimal for root growth, novel genes encoding cytokinin oxidases (which are cytokinin metabolizing enzymes) were cloned from *Arabidopsis thaliana* (designated *AtCKX*) and were subsequently expressed under a strong constitutive promoter in transgenic tobacco and *Arabidopsis*. Transformants showing *AtCKX* mRNA expression and increased cytokinin oxidase activity also manifested enhanced formation and growth of roots. Negative effects on shoot growth were also observed. The latter is in accordance with the constitutive expression of the cytokinin oxidase gene in these plants, illustrating the

importance of confined expression of the cytokinin oxidase gene for general plant growth properties. Containment of cytokinin oxidase activity can be achieved by using cell-, tissue- or organ-specific promoters, since cytokinin degradation is a process limited to the tissues or cells that express the CKX protein, this in contrast to approaches relying on hormone synthesis, as explained above.

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The observed negative effects of cytokinin oxidase expression on shoot growth demonstrate that cytokinin oxidases are interesting targets for the design of or screening for growth-promoting chemicals. Such chemicals should inhibit cytokinin oxidase activity, should preferably not be transported to the root and should be rapidly degraded in soil, so that application of these chemicals will not inhibit root growth. Cytokinins also delay leaf senescence, which means that positive effects will include both growth and maintenance of photosynthetic tissues. In addition, the observation that cytokinins delay senescence, enhance greening (chlorophyll content) of leaves and reduce shoot apical dominance shows that strategies based on suppressing CKX activity (such as antisense, ribozyme, and cosuppression technology) in the aerial parts of the plant could result in delayed senescence, enhanced leaf greening and increased branching.

Similarly, the observed positive effects of cytokinin oxidase expression on root growth demonstrate that cytokinin oxidases are interesting targets for the design of or screening for herbicides. Such herbicides should inhibit cytokinin oxidase activity, should preferably not be transported to the shoot, and should be soluble and relatively stable in a solvent that can be administered to the root through the soil.

These effects of cytokinin oxidase overexpression on plant development and architecture were hitherto unknown and, as a consequence, the presented invention and its embodiments could not be envisaged.

Preferable embodiments of the invention relate to the positive effect of cytokinin oxidase expression on plant growth and architecture, and in particular on root growth and architecture. The cytokinin oxidase gene family contains at least six members in *Arabidopsis* (see examples below) and we have shown that there are quantitative differences in the effects achieved with some of these genes in transgenic plants. It is anticipated that functional homologs of the described *Arabidopsis* cytokinin oxidases can be isolated from other organisms, given the evidence for the presence of cytokinin oxidase activity in many green plants (Hare and van Staden, Physiol Plant 91:128-136, 1994; Jones and Schreiber, Plant Growth Reg 23:123-134, 1997), as well as in other organisms (Armstrong, in Cytokinins: Chemistry, Activity and Function. Eds Mok and

Mok, CRC Press, pp139-154, 1994). Therefore, the sequence of the cytokinin oxidase, functional in the invention, need not to be identical to those described herein. It is envisaged that other genes with cytokinin oxidase activity or with any other cytokinin metabolizing activity (see Zažímalová *et al.*, Biochemistry and Molecular Biology of Plant Hormones, Hooykaas, Hall and Libbenga (Eds.), Elsevier Science, pp141-160, 1997) can also be used for the purpose of this invention. Similarly, genes encoding proteins that would increase endogenous cytokinin metabolizing activity can also be used for the purpose of this invention.

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According to a first embodiment, the present invention relates to a method for stimulating root growth and/or enhancing the formation of lateral and adventitious roots and/or altering root geotropism comprising expression of a plant cytokinin oxidase or comprising expression of another protein that reduces the level of active cytokinins in plants or plant parts.

In the context of the present invention it should be understood that the term "expression" relates to an "enhanced and/or ectopic expression" of a plant cytokinin oxidase or any other protein that reduces the level of active cytokinins in plants. It should be clear that herewith an enhanced expression of the plant cytokinin oxidase as well as "de novo" expression of plant cytokinin oxidases or of said other proteins is meant.

Alternatively, said other protein enhances the cytokinin metabolizing activity of a plant cytokinin oxidase.

According to a further embodiment, the present invention relates to a method for stimulating root growth and/or enhancing the formation of lateral and adventitious roots and/or altering root geotropism and/or increasing yield and/or enhancing early vigor and/or modifying root/shoot ratio and/or improving resistance to lodging and/or increasing drought tolerance and/or promoting in vitro propagation of explants, comprising expression of a plant cytokinin oxidase or comprising expression of another protein that reduces the level of active cytokinins in plants or plant parts.

According to a preferred embodiment, the present invention relates to a method for stimulating root growth resulting in an increase of root mass by overexpression of a cytokinin oxidase, preferably a cytokinin oxidase according to the invention, or another protein that reduces the level of active cytokinins in plants or plant parts, preferably in roots.

Higher root biomass production due to overexpression of growth promoting sequences has a direct effect on the yield and an indirect effect of production of compounds

produced by root cells or transgenic root cells or cell cultures of said transgenic root cells. One example of an interesting compound produced in root cultures is shikonin, the yield of which can be advantageously enhanced by said methods.

According to another embodiment, the present invention relates to an isolated nucleic acid encoding a novel plant protein having cytokinin oxidase activity or encoding an immunologically active and/or functional fragment of such a protein selected from the group consisting of:

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- (a) nucleic acid sequences comprising a DNA sequence as given in any of SEQ ID NOs 1, 3, 5, 7, 9 or 11, or the complement thereof,
- (b) nucleic acid sequences comprising the RNA sequences corresponding to any of SEQ ID NOs 1, 3, 5, 7, 9 or 11, or the complement thereof,
 - (c) nucleic acid sequences hybridizing to the nucleotide sequence as defined in (a) or (b) under stringent hybridization conditions,
 - (d) nucleic acid sequences encoding a protein with an amino acid sequence which is at least 35% similar, preferably 36%, 37%, 38%, 39% similar, more preferably 40%, 41%, 42%, 43%, 44% similar, most preferably 45%, 50%, 60%,70%, 80% 90% or 95% similar to the amino acid sequence as given in SEQ ID NO 2,
 - (e) nucleic acid sequences encoding a protein with an amino acid sequence which is at least 35% similar, preferably 36%, 37%, 38%, 39% similar, more preferably 40%, 41%, 42%, 43%, 44% similar, most preferably 45%, 50%, 60%,70%, 80% 90% or 95% similar to the amino acid sequence as given in SEQ ID NO 4,
 - (f) nucleic acid sequences encoding a protein with an amino acid sequence which is at least 35% similar, preferably 36%, 37%, 38%, 39% similar, more preferably 40%, 41%, 42%, 43%, 44% similar, most preferably 45% 50%, 60%,70%, 80% 90% or 95% similar to the amino acid sequence as given in SEQ ID NO 6,
 - (g) nucleic acid sequences encoding a protein with an amino acid sequence which is at least 35% similar, preferably 36%, 37%, 38%, 39% similar, more preferably 40%, 41%, 42%, 43%, 44% similar, most preferably 45% 50%, 60%,70%, 80% 90% or 95% similar to the amino acid sequence as given in SEQ ID NO 8,
- (h) nucleic acid sequences encoding a protein with an amino acid sequence which is at least 35% similar, preferably 36%, 37%, 38%, 39% similar, more preferably 40%, 41%, 42%, 43%, 44% similar, most preferably 45% 50%, 60%,70%, 80% 90% or 95% similar to the amino acid sequence as given in SEQ ID NO 10,

- (i) nucleic acid sequences encoding a protein with an amino acid sequence which is at least 35% similar, preferably 36%, 37%, 38%, 39% similar, more preferably 40%, 41%, 42%, 43%, 44% similar, most preferably 45% 50%, 60%,70%, 80% 90% or 95% similar to the amino acid sequence as given in SEQ ID NO 12,
- 5 (j) nucleic acid sequences encoding a protein comprising the amino acid sequence as given in any of SEQ ID NOs 2, 4, 6, 8, 10 or 12,

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- (k) nucleic acid sequences which are degenerated as a result of the genetic code to a nucleotide sequence of a nucleic acid as given in any of SEQ ID NOs 1, 3, 5, 7, 9 or 11, or as defined in (a) to (j),
- (I) nucleic acid sequences which are diverging due to the differences in codon usage between the organisms to a nucleotide sequence encoding a protein as given in any of SEQ ID NOs 2, 4, 6, 8, 10, 12 or as defined in (a) to (j),
- (m) nucleic acid sequences which are diverging due to the differences between alleles encoding a protein as given in SEQ ID NOs 2, 4, 6, 8, 10 or 12, or as defined in (a) to (j),
- (n) nucleic acid sequences encoding an immunologically active and/or functional fragment of a protein encoded by a DNA sequence as given in any of SEQ ID NOs 1, 3, 5, 7, 9 or 11, or a functional fragment of a nucleic acid as defined in any one of (a) to (m), and,
- (o) nucleic acid sequences encoding a protein as defined in SEQ ID NO 2, 4, 6, 8, 10 or 12 or a nucleic acid as defined in any one of (a) to (n) characterized in that said sequence is a cDNA sequence.

The invention also relates to an isolated nucleic acid according to claim 2 which is DNA, cDNA, genomic DNA or synthetic DNA, or RNA wherein T is replaced by U.

- The invention also relates to a nucleic acid molecule of at least 15 nucleotides in length hybridizing specifically with or specifically amplifying a nucleic acid of claim 2 or 3.
 - According to another embodiment, the invention also relates to a vector comprising a nucleic acid sequence according to claim 2 or 3. In a preferred embodiment, said vector is an expression vector wherein the nucleic acid sequence is operably linked to one or more control sequences allowing the expression of said sequence in prokaryotic and/or eukaryotic host cells.

It should be understood that for expression of the cytokinin oxidase genes of the invention in monocots, a nucleic acid sequence corresponding to the cDNA sequence should be used to avoid mis-splicing of introns in monocots.

The invention also relates to a host cell containing any of the nucleic acid molecules or vectors of the invention. Said host cell is chosen from the group coprising bacterial, insect, fungal, plant or animal cells.

Another embodiment of the invention relates to an isolated polypeptide encodable by a nucleic acid of claim 2 or 3, or a homologue or a derivative thereof, or an immunologically active and/or functional fragment thereof.

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In a preferred embodiment, the invention relates to a polypeptide according to claim 10 which has an amino acid sequence as given in SEQ ID NO 2, 4, 6, 8, 10 or 12, or a homologue or a derivative thereof, or an immunologically active and/or functional fragment thereof.

According to yet another embodiment, the invention relates to a method for producing a polypeptide according to claim 10 or 11 comprising culturing a host cell of claim 8 or 9 under conditions allowing the expression of the polypeptide and recovering the produced polypeptide from the culture.

The invention also relates to an antibody specifically recognizing a polypeptide of claim 10 or 11 or a specific epitope thereof.

The invention further relates to a method for the production of transgenic plants, plant cells or plant tissues comprising the introduction of a nucleic acid molecule according to any of claims 2 or 3 in an expressible format or a vector according to claim 6 or 7 in said plant, plant cell or plant tissue.

The invention also relates to a method for the production of altered plants, plant cells or plant tissues comprising the introduction of a polypeptide of claim 10 or 11 directly into a cell, a tissue or an organ of said plant.

According to another embodiment, the invention relates to a method for effecting the expression of a polypeptide of claim 10 or 11 comprising the introduction of a nucleic acid molecule of claim 2 or 3 operably linked to one or more control sequences or a vector according to claim 6 or 7 stably into the genome of a plant cell.

The invention further relates to a method of claim 15 or 16 further comprising regenerating a plant from said plant cell.

The invention also relates to a transgenic plant cell comprising a nucleic acid sequence of claim 2 or 3 which is operably linked to regulatory elements allowing transcription and/or expression of said nucleic acid in plant cells or obtainable by a method of claim 15 or 16.

According to a preferred embodiment, the invention relates to a transgenic plant cell of claim 18 wherein said nucleic acid of claim 2 or 3 is stably integrated into the genome of said plant cell.

The invention also relates to a transgenic plant or plant tissue comprising plant cells of claim 18 or 19 and to a harvestable part of said transgenic plant.

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According to the invention, the harvestable part of said transgenic plant is selected from the group consisting of seeds, leaves, fruits, stem cultures, roots, tubers, rhizomes and bulbs.

The invention also relates to the progeny derived from any of the plants or plant parts of any of claims 20 to 22.

According to another embodiment, the invention relates to a method for stimulating root growth comprising expression of a nucleic acid of claim 2 or 3 or comprising expression of another protein that reduces the level of active cytokinins in plants or plant parts.

The invention further relates to a method for enhancing the formation of lateral and adventitious roots comprising expression of a nucleic acid of claim 2 or 3 or comprising expression of another protein that reduces the level of active cytokinins in plants or plant parts.

The invention also relates to method for altering root geotropism comprising altering the expression of a nucleic acid of claim 2 or 3 or comprising expression of another protein that that reduces the level of active cytokinins in plants or plant parts.

The invention also relates to methods for enhancing early vigor and/or for modifying root/shoot ratio and/or for improving resistance to lodging and/or for increasing drought tolerance and/or for promoting *in vitro* propagation of explants comprising expression of a nucleic acid of claim 2 or 3 or comprising expression of another protein that reduces the level of active cytokinins in plants or plant parts.

According to a preferred embodiment the invention relates to a method for delaying leaf senescence comprising downregulation of expression of any of the cytokinin oxidases of the inventiong in leaves, preferably in senescing leaves. Also the invention relates to a method for altering leaf senescence comprising expression of one of the cytokinin oxidases in senescing leaves.

The invention further relates to methods for increasing the root size or the size of the root meristem comprising expression of a nucleic acid of claim 2 or 3 or comprising expression of another protein that reduces the level of active cytokinins in plants or plant parts, preferably in roots.

According to yet another embodiment, the invention relates to a method for increasing the size of the shoot meristem comprising downregulation of expression of a nucleic acid of claim 2 or 3, preferably in shoots.

The invention also relates to methods for increasing leaf thickness comprising expression of a nucleic acid of claim 2 or 3 or comprising expression of another protein that reduces the level of active cytokinins in plants or plant parts, preferably in leaves.

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The invention also relates to a method for reducing the vessel size comprising expression of a nucleic acid of claim 2 or 3 or comprising expression of another protein that reduces the level of active cytokinins in vessels.

The invention further relates to a method for increasing the vessel size comprising downregulation of expression of a nucleic acid of claim 2 or 3 in vessels.

According to another embodiment, the invention relates to a method for improving standability of seedlings comprising expression of a nucleic acid of claim 2 or 3 or comprising expression of another protein that reduces the level of active cytokinins in seedlings.

Furthermore, the invention relates to any of the above described methods, preferably a method of any of claims 24 to 26, said method leading to an increase in yield.

The invention further relates to any of the above described methods, preferably a method of any of claims 24 to 27, wherein said expression of said nucleic acid occurs under the control of a strong constitutive promoter.

In a preferred embodiment the invention relates to any of the above described methods, preferably a method of any of claims 24 to 27, wherein said expression of said nucleic acid occurs under the control of a promoter that is preferentially expressed in roots.

According to yet another embodiment, the invention relates to a method for modifying cell fate and/or modifying plant development and/or modifying plant morphology and/or modifying plant biochemistry and/or modifying plant physiology and/or modifying the cell cycle progression rate comprising the modification of expression in particular cells, tissues or organs of a plant, of a nucleic acid sequence of claim 2 or 3.

The invention also relates to a method for obtaining enhanced growth, and/or increased yield and/or altered senescence of a plant cell, tissue and/or organ and/or increased frequence of formation of lateral organs in a plant, comprising the ectopic expression of a nucleic acid sequence of claim 2 or 3.

The invention also relates to a method for promoting and extending cell division activity in cells in adverse growth conditions and/or in stress, comprising the ectopic expression of a nucleic acid sequence of claim 2 or 3.

According to yet another embodiment, the invention relates to a method for identifying and obtaining proteins interacting with a polypeptide of claim 10 or 11 comprising a screening assay wherein a polypeptide of claim 10 or 11 is used.

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In a more preferred embodiment, the invention relates to a method for identifying and obtaining proteins interacting with a polypeptide of claim 10 or 11 comprising a two-hybrid screening assay wherein a polypeptide of claim 10 or 11 as a bait and a cDNA library as prey are used.

The invention further relates to a method for modulating the interaction between a polypeptide of claim 10 or 11 and interacting protein partners obtainable by a method according to claim 30 or 31.

In a further embodiment, the invention relates to a method for identifying and obtaining compounds interacting with a polypeptide of claim 10 or 11 comprising the steps of:

- a) providing a two-hybrid system wherein a polypeptide of claim 10 or 11 and an interacting protein partner obtainable by a method according to claim 30 or 31 are expressed,
- b) interacting said compound with the complex formed by the expressed polypeptides as defined in a), and,
- c) performing (real-time) measurement of interaction of said compound with said polypeptide or the complex formed by the expressed polypeptides as defined in a).

The invention further relates to a method for identifying compounds or mixtures of compounds which specifically bind to a polypeptide of claim 10 or 11, comprising:

- a) combining a polypeptide of claim 10 or 11 with said compound or mixtures of compounds under conditions suitable to allow complex formation, and,
- b) detecting complex formation, wherein the presence of a complex identifies a compound or mixture which specifically binds said polypeptide.
- The invention also relates to a method of any of claims 30 to 34 wherein said compound or mixture inhibits the activity of said polypeptide of claim 10 or 11 and can be used for the rational design of chemicals.

According to another embodiment, the invention relates to the use of a compound or mixture identified by means of a method of any of claims 30 to 34 as a plant growth regulator or herbicide.

The invention also relates to a method for production of a plant growth regulator or herbicide composition comprising the steps of the method of any of claims 30 to 34 and formulating the compounds obtained from said steps in a suitable form for the application in agriculture or plant cell or tissue culture.

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The invention also relates to a method for increasing branching comprising expression of a nucleic acid of claim 2 or 3 in plants or plant parts, preferably in stems or axillary buds.

The invention also relates to a method for improving lodging resistance comprising expression of a nucleic acid of claim 2 or 3 in plants or plant parts, preferably in stems or axillary buds.

The invention also relates to a method for the design of or screening for growth-promoting chemicals or herbicides comprising the use of a nucleic acid of claim 2 or 3 or a vector of claim 6 or 7.

According to another embodiment, the invention relates to the use of a nucleic acid molecule of claim 2 or 3, the vector of claim 6 or 7, a polypeptide of claim 10 or 11 for increasing yield.

The invention also relates to the use of a nucleic acid molecule of claim 2 or 3, the vector of claim 6 or 7, a polypeptide of claim 10 or 11 for stimulating root growth.

The invention also relates to the use of a nucleic acid molecule of claim 2 or 3, the vector of claim 6 or 7, a polypeptide of claim 10 or 11 for enhancing the formation of lateral and adventitious roots.

The invention also relates to the use of a nucleic acid molecule of claim 2 or 3, the vector of claim 6 or 7, a polypeptide of claim 10 or 11 for altering root geotropism.

The invention further relates to the use of a nucleic acid molecule of claim 2 or 3, the vector of claim 6 or 7, a polypeptide of claim 10 or 11 for enhancing early vigor and/or for modifying root/shoot ratio and/or for improving resistance to lodging and/or for increasing drought tolerance and/or for promoting *in vitro* propagation of explants.

The invention also relates to the use of a nucleic acid molecule of claim 2 or 3, the recombinant vector of claim 6 or 7, a polypeptide of claim 10 or 11 for modifying plant development and/or for modifying plant morphology and/or for modifying plant biochemistry and/or for modifying plant physiology.

According to yet another embodiment, the invention relates to a diagnostic composition comprising at least a nucleic acid molecule of any of claims 2 to 5, the vector of claim 6 or 7, a polypeptide of claim 10 or 11 or an antibody of claim 13.

As mentioned supra, auxins and cytokinins act as antagonists in certain biological processes. For example, the cytokinin/auxin ratio regulates the production of roots and shoots with a high concentration of auxin resulting in organized roots and a high concentration of cytokinins resulting in shoot production. Another example is apical dominance, which is due to the inhibitory influence of auxins on lateral bud development. Treatment of lateral buds with cytokinins often causes the bud to grow, thus counteracting the auxin effect and modifying apical dominance. As disclosed in this invention, expression of cytokinin oxidases in tobacco and Arabidopsis results in enhanced root development and axillary shoot formation consistent with enhanced auxin effects. Auxins are also involved in the development of fruit. Treatment of female flower parts with auxin results in the development of parthenocarpic fruit in some plant species. Parthenocarpic fruit development has been genetically engineered in several horticultural crop plants through increased biosynthesis of auxins in the female reproductive organs (WO0105985).

Therefore, according to another embodiment, this invention relates to a method for inducing the parthenocarpic trait in plants, said method consisting of downregulating the expression of one or more cytokinin oxidases or of another protein that reduces the level of active cytokinins in plants or plant parts, preferably in the female reproductive organs such as the placenta, ovules and tissues derived therefrom. The DefH9 promoter region from Antirrhinum majus or one of its homologues which confers a high expression specificity in placenta and ovules can be used for this purpose.

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DEFINITIONS AND ELABORATIONS TO THE EMBODIMENTS

Those skilled in the art will be aware that the invention described herein is subject to variations and modifications other than those specifically described. It is to be understood that the invention described herein includes all such variations and modifications. The invention also includes all such steps, features, compositions and compounds referred to or indicated in this specification, individually or collectively, and any and all combinations of any or more of said steps or features.

The present invention is applicable to any plant, in particular a monocotyledonous plants and dicotyledonous plants including a fodder or forage legume, ornamental plant, food

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crop, tree, or shrub selected from the list comprising Acacia spp., Acer spp., Actinidia spp., Aesculus spp., Agathis australis, Albizia amara, Alsophila tricolor, Andropogon spp., Arachis spp, Areca catechu, Astelia fragrans, Astragalus cicer, Baikiaea plurijuga, Betula spp., Brassica spp., Bruguiera gymnorrhiza, Burkea africana, Butea frondosa, Cadaba farinosa, Calliandra spp, Camellia sinensis, Canna indica, Capsicum spp., Cassia spp., Centroema pubescens, Chaenomeles spp., Cinnamomum cassia, Coffea arabica, Colophospermum mopane, Coronillia varia, Cotoneaster serotina, Crataegus spp., Cucumis spp., Cupressus spp., Cyathea dealbata, Cydonia oblonga, Cryptomeria japonica, Cymbopogon spp., Cynthea dealbata, Cydonia oblonga, Dalbergia monetaria, Davallia divaricata, Desmodium spp., Dicksonia squarosa, Diheteropogon amplectens, Dioclea spp, Dolichos spp., Dorycnium rectum, Echinochloa pyramidalis, Ehrartia spp., Eleusine coracana, Eragrestis spp., Erythrina spp., Eucalyptus spp., Euclea schimperi, Eulalia villosa, Fagopyrum spp., Feijoa sellowiana, Fragaria spp., Flemingia spp, Freycinetia banksii, Geranium thunbergii, Ginkgo biloba, Glycine javanica, Gliricidia spp, Gossypium hirsutum, Grevillea spp., Guibourtia coleosperma, Hedysarum spp., Hemarthia altissima, Heteropogon contortus, Hordeum vulgare, Hyparrhenia rufa, Hypericum erectum, Hyperthelia dissoluta, Indigo incarnata, Iris spp., Leptarrhena pyrolifolia, Lespediza spp., Lettuca spp., Leucaena leucocephala, Loudetia simplex, Lotonus bainesii, Lotus spp., Macrotyloma axillare, Malus spp., Manihot esculenta, Medicago sativa, Metasequoia glyptostroboides, Musa sapientum, Nicotianum spp., Onobrychis spp., Ornithopus spp., Oryza spp., Peltophorum africanum, Pennisetum spp., Persea gratissima, Petunia spp., Phaseolus spp., Phoenix canariensis, Phormium cookianum, Photinia spp., Picea glauca, Pinus spp., Pisum sativum, Podocarpus totara, Pogonarthria fleckii, Pogonarthria squarrosa, Populus spp., Prosopis cineraria, Pseudotsuga menziesii, Pterolobium stellatum, Pyrus communis, Quercus spp., Rhaphiolepsis umbellata, Rhopalostylis sapida, Rhus natalensis, Ribes grossularia, Ribes spp., Robinia pseudoacacia, Rosa spp., Rubus spp., Salix spp., Schyzachyrium Sciadopitys verticillata, sanguineum, Sequoia sempervirens, Sequoiadendron giganteum, Sorghum bicolor, Spinacia spp., Sporobolus fimbriatus, Stiburus alopecuroides, Stylosanthos humilis, Tadehagi spp, Taxodium distichum, Themeda triandra, Trifolium spp., Triticum spp., Tsuga heterophylla, Vaccinium spp., Vicia spp. Vitis vinifera, Watsonia pyramidata, Zantedeschia aethiopica, Zea mays, amaranth, artichoke, asparagus, broccoli, brussel sprout, cabbage, canola, carrot, cauliflower, celery, collard greens, flax, kale, lentil, oilseed rape, okra, onion, potato, rice, soybean,

straw, sugarbeet, sugar cane, sunflower, tomato, squash, and tea, amongst others, or the seeds of any plant specifically named above or a tissue, cell or organ culture of any of the above species.

Throughout this specification, unless the context requires otherwise the word "comprise", and variations such as "comprises" and "comprising", will be understood to imply the inclusion of a stated integer or step or group of integers or steps but not the exclusion of any other integer or step or group of integers or steps.

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As used herein, the term "derived from" shall be taken to indicate that a particular integer or group of integers has originated from the species specified, but has not necessarily been obtained directly from the specified source.

The terms "protein(s)", "peptide(s)" or "oligopeptide(s)", when used herein refer to amino acids in a polymeric form of any length. Said terms also include known amino acid modifications such as disulphide bond formation, cysteinylation, oxidation, glutathionylation, methylation, acetylation, farnesylation, biotinylation, stearoylation, formylation, lipoic acid addition, phosphorylation, sulphation, ubiquitination, myristoylation, palmitoylation, geranylgeranylation, cyclization (e.g. pyroglutamic acid oxidation, deamidation, dehydration, glycosylation formation), (e.g. hexosamines, N-acetylhexosamines, deoxyhexoses, hexoses, sialic acid etc.) and acylation as well as non-naturally occurring amino acid residues, L-amino acid residues and D-amino acid residues.

"Homologues" of a protein of the invention are those peptides, oligopeptides, polypeptides, proteins and enzymes which contain amino acid substitutions, deletions and/or additions relative to the said protein with respect to which they are a homologue, without altering one or more of its functional properties, in particular without reducing the activity of the resulting. For example, a homologue of said protein will consist of a bioactive amino acid sequence variant of said protein. To produce such homologues, amino acids present in the said protein can be replaced by other amino acids having similar properties, for example hydrophobicity, hydrophilicity, hydrophobic moment, antigenicity, propensity to form or break α -helical structures or β -sheet structures, and so on. An overview of physical and chemical properties of amino acids is given in Table 1.

Substitutional variants of a protein of the invention are those in which at least one residue in said protein amino acid sequence has been removed and a different residue inserted in its place. Amino acid substitutions are typically of single residues, but may be clustered depending upon functional constraints placed upon the polypeptide; insertions

will usually be of the order of about 1-10 amino acid residues. and deletions will range from about 1-20 residues. Preferably, amino acid substitutions will comprise conservative amino acid substitutions, such as those described *supra*.

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Table 1. Properties of naturally occurring amino acids.

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Charge properties / hydrophobicity	Side group	Amino Acid
nonpolar	Aliphatic	ala, ile, leu, val
hydrophobic	aliphatic, S-containing	met
	aromatic	phe, trp
	imino	pro
polar uncharged	Aliphatic	gly
	amide	asn, gln
	aromatic	tyr
	hydroxyl	ser, thr
	sulfhydryl	cys
positively charged	Basic	arg, his, lys
negatively charged	Acidic	asp, glu

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Insertional amino acid sequence variants of a protein of the invention are those in which one or more amino acid residues are introduced into a predetermined site in said protein. Insertions can comprise amino-terminal and/or carboxy-terminal fusions as well as intra-sequence insertions of single or multiple amino acids. Generally, insertions within the amino acid sequence will be smaller than amino or carboxyl terminal fusions, of the order of about 1 to 10 residues. Examples of amino- or carboxy-terminal fusion proteins or peptides include the binding domain or activation domain of a transcriptional activator as used in a two-hybrid system, phage coat proteins, (histidine)₆-tag, glutathione S-transferase, protein A, maltose-binding protein, dihydrofolate reductase, Tag•100 epitope (EETARFQPGYRS), c-myc epitope (EQKLISEEDL), FLAG®-epitope

(DYKDDDK), lacZ, CMP (calmodulin-binding peptide), HA epitope (YPYDVPDYA), protein C epitope (EDQVDPRLIDGK) and VSV epitope (YTDIEMNRLGK).

Deletional variants of a protein of the invention are characterised by the removal of one or more amino acids from the amino acid sequence of said protein.

Amino acid variants of a protein of the invention may readily be made using peptide synthetic techniques well known in the art, such as solid phase peptide synthesis and the like, or by recombinant DNA manipulations. The manipulation of DNA sequences to produce variant proteins which manifest as substitutional, insertional or deletional variants are well known in the art. For example, techniques for making substitution mutations at predetermined sites in DNA having known sequence are well known to those skilled in the art, such as by M13 mutagenesis, T7-Gen in vitro mutagenesis kit (USB, Cleveland, OH), QuickChange Site Directed mutagenesis kit (Stratagene, San Diego, CA), PCR-mediated site-directed mutagenesis or other site-directed mutagenesis protocols.

In the current invention "homology" and "similarity" percentages between proteins are calculated using computer programs known in the art such as the Dnastar/MegAlign programs in combination with the Clustal method.

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"Derivatives" of a protein of the invention are those peptides, oligopeptides, polypeptides, proteins and enzymes which comprise at least about five contiguous amino acid residues of said polypeptide but which retain the biological activity of said protein. A "derivative" may further comprise additional naturally-occurring, altered glycosylated, acylated or non-naturally occurring amino acid residues compared to the amino acid sequence of a naturally-occurring form of said polypeptide. Alternatively or in addition, a derivative may comprise one or more non-amino acid substituents compared to the amino acid sequence of a naturally-occurring form of said polypeptide, for example a reporter molecule or other ligand, covalently or non-covalently bound to the amino acid sequence such as, for example, a reporter molecule which is bound thereto to facilitate its detection.

With "immunologically active" is meant that a molecule or specific fragments thereof such as specific epitopes or haptens are recognized by, i.e. bind to antibodies. Specific epitopes may be determined using, for example, peptide scanning techniques as described in Geysen *et al.* (1996) (Geysen, H.M., Rodda, S.J. and Mason, T.J. (1986). A priori delineation of a peptide which mimics a discontinuous antigenic determinant. *Mol. Immunol.* 23, 709-715.).

The term "fragment of a sequence" or "part of a sequence" means a truncated sequence of the original sequence referred to. The truncated sequence (nucleic acid or protein sequence) can vary widely in length; the minimum size being a sequence of sufficient size to provide a sequence with at least a comparable function and/or activity or the original sequence referred to (e. g. "functional fragment"), while the maximum size is not critical. In some applications, the maximum size usually is not substantially greater than that required to provide the desired activity and/or function(s) of the original sequence. Typically, the truncated amino acid sequence will range from about 5 to about 60 amino acids in length. More typically, however, the sequence will be a maximum of about 50 amino acids in length, preferably a maximum of about 60 amino acids. It is usually desirable to select sequences of at least about 10, 12 or 15 amino acids, up to a maximum of about 20 or 25 amino acids.

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Functional fragments include those comprising an epitope which is specific for the proteins according to the invention. Preferred functional fragments have a length of at least, for example, 5, 10, 25, 100, 150 or 200 amino acids.

In the context of the current invention are embodied homologues, derivatives and/or immunologically active and/or functional fragments of inventive cytokinin oxidases as defined supra. Particularly preferred homologues, derivatives and/or immunologically active and/or functional fragments of the cytokinin oxidase proteins which are contemplated for use in the current invention are derived from plants, more specifically from *Arabidopsis thaliana*, even more specifically said cytokinin oxidases are the *Arabidopsis thaliana* (At)CKX, or are capable of being expressed therein. The present invention clearly contemplates the use of functional homologues or derivatives and/or immunologically active fragments of the AtCKX proteins and is not to be limited in application to the use of a nucleotide sequence encoding one of said AtCKX proteins.

Any of said proteins, polypeptides, peptides and fragments thereof can be produced in a biological system, e.g. a cell culture. Alternatively any of said proteins, polypeptides, peptides and fragments thereof can be chemically manufactured e.g. by solid phase peptide synthesis. Said proteins or fragments thereof can be part of a fusion protein as is the case in e.g. a two-hybrid assay which enables e.g. the identification of proteins interacting with a cytokinin oxidase according to the invention.

The proteins or fragments thereof are furthermore useful e.g. to modulate the interaction between a cytokinin oxidase according to the invention and interacting protein partners

obtained by a method of the invention. Chemically synthesized peptides are particularly useful e.g. as a source of antigens for the production of antisera and/or antibodies.

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"Antibodies" include monoclonal, polyclonal, synthetic or heavy chain camel antibodies as well as fragments of antibodies such as Fab, Fv or scFv fragments. Monoclonal antibodies can be prepared by the techniques as described in e.g. Liddle and Cryer (1991) which comprise the fusion of mouse myeloma cells to spleen cells derived from immunized animals. Furthermore, antibodies or fragments thereof to a molecule or fragments thereof can be obtained by using methods as described in e.g. Harlow and Lane (1988). In the case of antibodies directed against small peptides such as fragments of a protein of the invention, said peptides are generally coupled to a carrier protein before immunization of animals. Such protein carriers include keyhole limpet hemocyanin (KLH), bovine serum albumin (BSA), ovalbumin and Tetanus toxoid. The carrier protein enhances the immune response of the animal and provides epitopes for T-cell receptor binding sites. The term "antibodies" furthermore includes derivatives thereof such as labelled antibodies. Antibody labels include alkaline phosphatase, PKH2, PKH26, PKH67, fluorescein (FITC), Hoechst 33258, R-phycoerythrin (PE), rhodamine (TRITC), Quantum Red, Texas Red, Cy3, biotin, agarose, peroxidase and gold spheres. Tools in molecular biology relying on antibodies against a protein include protein gel blot analysis, screening of expression libraries allowing gene identification, protein quantitative methods including ELISA and RIA, immunoaffinity purification of proteins, immunoprecipitation of proteins (see e.g. Example 5) and immunolocalization (see e.g. Example 6). Other uses of antibodies and especially of peptide antibodies include the study of proteolytic processing (Loffler et al. 1994, Woulfe et al. 1994), determination of protein active sites (Lerner 1982), the study of precursor and posttranslational processing (Baron and Baltimore 1982, Lerner et al. 1981, Semier et al. 1982), identification of protein domains involved in protein-protein interactions (Murakami et al. 1992) and the study of exon usage in gene expression (Tamura et al. 1991).

Embodied in the current invention are antibodies specifically recognizing a cytokinin oxidase or homologue, derivative or fragment thereof as defined supra. Preferably said cytokinin oxidase is a plant cytokinin oxidase, more specifically one of the *Arabidopsis thaliana* cytokinin oxidases (AtCKX).

The terms "gene(s)", "polynucleotide(s)", "nucleic acid sequence(s)", "nucleotide sequence(s)", or "nucleic acid molecule(s)", when used herein refer to nucleotides, either

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ribonucleotides or deoxyribonucleotides or a combination of both, in a polymeric form of any length. Said terms furthermore include double-stranded and single-stranded DNA and RNA. Said terms also include known nucleotide modifications such as methylation, cyclization and 'caps' and substitution of one or more of the naturally occurring nucleotides with an analog such as inosine. Modifications of nucleotides include the addition of acridine, amine, biotin, cascade blue, cholesterol, Cy3[®], Cy5[®], Cy5.5[®] Dabcyl, digoxigenin, dinitrophenyl, Edans, 6-FAM, fluorescein, 3'-glyceryl, HEX, IRD-700, IRD-800, JOE, phosphate psoralen, rhodamine, ROX, thiol (SH), spacers, TAMRA, TET, AMCA-S®, SE, BODIPY®, Marina Blue®, Pacific Blue®, Oregon Green®, Rhodamine Green®, Rhodamine Red®, Rhodol Green® and Texas Red®. Polynucleotide backbone modifications methylphosphonate, include 2'-OMe-methylphosphonate RNA, phosphorothiorate, RNA, 2'-OMeRNA. Base modifications include 2-amino-dA, 2aminopurine, 3'-(ddA), 3'dA(cordycepin), 7-deaza-dA, 8-Br-dA, 8-oxo-dA, N⁶-Me-dA, abasic site (dSpacer), biotin dT, 2'-OMe-5Me-C, 2'-OMe-propynyl-C, 3'-(5-Me-dC), 3'-(ddC), 5-Br-dC, 5-I-dC, 5-Me-dC, 5-F-dC, carboxy-dT, convertible dA, convertible dC, convertible dG, convertible dT, convertible dU, 7-deaza-dG, 8-Br-dG, 8-oxo-dG, O⁶-MedG, S6-DNP-dG, 4-methyl-indole, 5-nitroindole, 2'-OMe-inosine, 2'-dl, 06-phenyl-dl, 4methyl-indole, 2'-deoxynebularine, 5-nitroindole, 2-aminopurine, dP(purine analogue), dK(pyrimidine analogue), 3-nitropyrrole, 2-thio-dT, 4-thio-dT, biotin-dT, carboxy-dT, O4-Me-dT, O⁴-triazol dT, 2'-OMe-propynyl-U, 5-Br-dU, 2'-dU, 5-F-dU, 5-I-dU, O⁴-triazol dU. Said terms also encompass peptide nucleic acids (PNAs), a DNA analogue in which the backbone is a pseudopeptide consisting of N-(2-aminoethyl)-glycine units rather than a sugar. PNAs mimic the behaviour of DNA and bind complementary nucleic acid strands. The neutral backbone of PNA results in stronger binding and greater specificity than normally achieved. In addition, the unique chemical, physical and biological properties of PNA have been exploited to produce powerful biomolecular tools, antisense and antigene agents, molecular probes and biosensors.

The present invention also advantageously provides nucleic acid sequences of at least approximately 15 contiguous nucleotides of a nucleic acid according to the invention and preferably from 15 to 50 nucleotides. These sequences may, advantageously be used as probes to specifically hybridise to sequences of the invention as defined above or primers to initiate specific amplification or replication of sequences of the invention as defined above, or the like. Such nucleic acid sequences may be produced according to techniques well known in the art, such as by recombinant or synthetic means. They may

also be used in diagnostic kits or the like for detecting the presence of a nucleic acid according to the invention. These tests generally comprise contacting the probe with the sample under hybridising conditions and detecting the presence of any duplex or triplex formation between the probe and any nucleic acid in the sample.

Advantageously, the nucleic acid sequences, according to the invention may be produced using such recombinant or synthetic means, such as for example using PCR cloning mechanisms which generally involve making a pair of primers, which may be from approximately 15 to 50 nucleotides to a region of the gene which is desired to be cloned, bringing the primers into contact with mRNA, cDNA or genomic DNA from a cell, performing a polymerase chain reaction under conditions which bring about amplification of the desired region, isolating the amplified region or fragment and recovering the amplified DNA. Generally, such techniques as defined herein are well known in the art, such as described in Sambrook et al. (Molecular Cloning: a Laboratory Manual, 1989).

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A "coding sequence" or "open reading frame" or "ORF" is defined as a nucleotide sequence that can be transcribed into mRNA and/or translated into a polypeptide when placed under the control of appropriate control sequences or regulatory sequences, i.e. when said coding sequence or ORF is present in an expressible format. Said coding sequence of ORF is bounded by a 5' translation start codon and a 3' translation stop codon. A coding sequence or ORF can include, but is not limited to RNA, mRNA, cDNA, recombinant nucleotide sequences, synthetically manufactured nucleotide sequences or genomic DNA. Said coding sequence or ORF can be interrupted by intervening nucleic acid sequences.

Genes and coding sequences essentially encoding the same protein but isolated from different sources can consist of substantially divergent nucleic acid sequences. Reciprocally, substantially divergent nucleic acid sequences can be designed to effect expression of essentially the same protein. Said nucleic acid sequences are the result of e.g. the existence of different alleles of a given gene, of the degeneracy of the genetic code or of differences in codon usage. Thus, as indicated in Table 2, amino acids such as methionine and tryptophan are encoded by a single codon whereas other amino acids such as arginine, leucine and serine can each be translated from up to six different codons. Differences in preferred codon usage are illustrated in Table 3 for Agrobacterium tumefaciens (a bacterium), A. thaliana, M. sativa (two dicotyledonous plants) and Oryza sativa (a monocotyledonous plant). To extract one example, the codon GGC (for glycine) is the most frequently used codon in A. tumefaciens (36.2 ‰),

is the second most frequently used codon in *O. sativa* but is used at much lower frequencies in *A. thaliana* and *M. sativa* (9 ‰ and 8.4 ‰, respectively). Of the four possible codons encoding glycine (see Table 2), said GGC codon is most preferably used in *A. tumefaciens* and *O. sativa*. However, in *A. thaliana* this is the GGA (and GGU) codon whereas in *M. sativa* this is the GGU (and GGA) codon.

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DNA sequences as defined in the current invention can be interrupted by intervening sequences. With "intervening sequences" is meant any nucleic acid sequence which disrupts a coding sequence comprising said inventive DNA sequence or which disrupts the expressible format of a DNA sequence comprising said inventive DNA sequence. Removal of the intervening sequence restores said coding sequence or said expressible format. Examples of intervening sequences include introns and mobilizable DNA sequences such as transposons. With "mobilizable DNA sequence" is meant any DNA sequence that can be mobilized as the result of a recombination event.

Table 2. Degeneracy of the genetic code.

Amino Acid	Three -letter code	One- letter code	Possible codons					
Alanine	Ala	Α	GCA	GCC	GCG	GCU		
Arginine	Arg	R	AGA	AGG	CGA	CGC	CGG	CGU
Asparagine	Asn	N	AAC	AAU				
Aspartic Acid	Asp	D	GAC	GAU				
Cysteine	Cys	C	UGC	UGU				
Glutamic Acid	Glu	E	GAA	GAG				
Glutamine	Gln	Q	CAA	CAG				
Glycine	Gly	G	GGA	GGC	GGG	GGU		
Histidine	His	Н	CAC	CAU				
Isoleucine	lle	1	AUA	AUC	AUU			
Leucine	Leu	L	UUA	UUG	CUA	CUC	CUG	CUU
Lysine	Lys	K	AAA	AAG				
Methionine	Met	M	AUG					
Phenylalanine	Phe	F	UUC	UUU				
Proline	Pro	ρ.	CCA	CCC	CCG	CCU		
Serine	Ser	S	AGC	AGU	UCA	UCC	UCG	UCU
Threonine	Thr	T	ACA	ACC	ACG	ACU		
Tryptophan	Trp	W	UGG					
Tyrosine	Tyr	Y	UAC	UAU				
Valine	Val	V	GUA	GUC	GUG	GUU		
			Possible "STOP" codons					
			UAA	UAG	UGA			

Table 3. Usage of the indicated codons in the different organisms given as frequency per thousand codons (http://www.kazusa.or.jp/codon).

Codon	Agrobacterium	Arabidopsis	Modioogo	0,000	
	tumefaciens	thaliana	Medicago	Oryza	
UUU	13.9	22.5	sativa	sativa	
UUC	24.3	20.7	24.1	11.3	
UUA	3.5	12.9	16.9	26.3	
UUG	13.2	21.0	10.4	4.7	
UCU	7.0	24.6	22.4	11.8	
UCC	14.8	10.8	19.8 7.7	10.1	
UCA	7.4	17.8	17.2	16.9	
UCG	18.2	8.9	3.2	9.7	
UAU	12.3	15.2	16.6	10.8	
UAC	10.3	13.7	14.0	9.2 20.6	
UAA	0.9	0.9	1.2	0.9	
UAG	0.6	0.5	0.8	0.8	
UGU	3.0	10.8	10.6	5.0	
UGC	7.4	7.2	5.8	14.3	
UGA	1.8	1.0	0.8	1.3	
UGG	12.2	12.7	10.0	12.8	
CUU	19.1	24.3	28.3	14.6	
CUC	25.7	15.9	12.0	28.0	
CUA	5.2	10.0	8.8	5.7	
CUG	31.6	9.9	8.5	22.1	
CCU	7.7	18.3	23.2	11.8	
CCC	10.6	5.3	5.3	12.5	
CCA	8.9	16.1	22.6	12.2	
CCG	20.7	8.3	3.6	16.7	
CAU	10.6	14.0	14.6	9.2	
CAC	9.1	8.7	9.1	14.6	
CAA	11.2	19.7	23.2	11.9	
CAG	24.9	15.2	12.3	24.6	
CGU	12.2	8.9	10.1	6.8	
CGC	25.5	3.7	4.2	15.9	
CGA	8.2	6.2	4.2	4.2	
CGG	13.2	4.8	1.8	9.7	
AUU	15.4	22.0	29.4	13.8	
AUC	36.9	18.5	14.7	25.5	
AUA	6.2	12.9	11.7	7.2	
AUG	24.7	24.5	21.7	24.4	
ACU	6.4	17.8	20.8	10.3	
ACC	20.9	10.3	11.7	18.6	
ACA	9.1	15.9	18.9	10.0	

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ACG	18.8	7.6	2.8	10.8
AAU	13.5	22.7	25.0	12.9
AAC	18.7	20.9	18.7	25.1
AAA	13.6	31.0	32.2	12.0
AAG	24.4	32.6	35.1	39.4
AGU	5.7	14.0	12.6	7.3
AGC	15.8	11.1	8.8	16.9
AGA	5.3	18.7	13.6	7.7
AGG	6.5	10.9	11.7	14.9
GUU	16.6	27.3	34.7	15.0
GUC	29.3	12.7	9.9	22.8
GUA	6.1	10.1	10.0	5.7
GUG	19.7	17.5	16.5	25.0
GCU	17.4	28.0	34.6	19.8
GCC	35.8	10.3	11.4	33.2
GCA	19.5	17.6	25.9	15.6
GCG	31.7	8.8	3.4	25.3
GAU	25.8	36.8	40.0	21.5
GAC	28.0	17.3	15.5	31.6
GAA	29.9	34.4	35.9	17.1
GAG	26.3	32.2	27.4	41.1
GGU	16.5	22.2	28.7	16.3
GGC	36.2	9.0	8.4	34.7
GGA	12.5	23.9	27.3	15.0
GGG	11.3	10.2	7.4	16.6

"Hybridization" is the process wherein substantially homologous complementary nucleotide sequences anneal to each other. The hybridization process can occur entirely in solution, i.e. both complementary nucleic acids are in solution. Tools in molecular biology relying on such a process include PCR, subtractive hybridization and DNA sequence determination. The hybridization process can also occur with one of the complementary nucleic acids immobilized to a matrix such as magnetic beads, Sepharose beads or any other resin. Tools in molecular biology relying on such a process include the isolation of poly (A+) mRNA. The hybridization process can furthermore occur with one of the complementary nucleic acids immobilized to a solid support such as a nitrocellulose or nylon membrane or immobilized by e.g. photolitography to e.g. a silicious glass support (the latter known as nucleic acid arrays or microarrays or as nucleic acid chips). Tools in molecular biology relying on such a process include RNA and DNA gel blot analysis, colony hybridization, plaque hybridization and microarray hybridization. In order to allow hybridization to occur, the nucleic acid molecules are generally thermally or chemically (e.g. by NaOH) denatured to melt a double strand into two single strands and/or to remove hairpins or other

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secondary structures from single stranded nucleic acids. The stringency of hybridization is influenced by conditions such as temperature, salt concentration and hybridization buffer composition. High stringency conditions for hybridization include high temperature and/or low salt concentration (salts include NaCl and Na3-citrate) and/or the inclusion of formamide in the hybridization buffer and/or lowering the concentration of compounds such as SDS (detergent) in the hybridization buffer and/or exclusion of compounds such as dextran sulfate or polyethylene glycol (promoting molecular crowding) from the hybridization buffer. Conventional hybridization conditions are described in e.g. Sambrook et al. (1989) but the skilled craftsman will appreciate that numerous different hybridization conditions can be designed in function of the known or the expected homology and/or length of the nucleic acid sequence. Sufficiently low stringency hybridization conditions are particularly preferred to isolate nucleic acids heterologous to the DNA sequences of the invention defined supra. Elements contributing to said heterology include allelism, degeneration of the genetic code and differences in preferred codon usage as discussed supra.

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Clearly, the current invention embodies the use of the inventive DNA sequences encoding a cytokinin oxidase, homologue, derivative or immunologically active and/or functional fragment thereof as defined higher in any method of hybridization. The current invention furthermore also related to DNA sequences hybridizing to said inventive DNA sequences. Preferably said cytokinin oxidase is a plant cytokinin oxidase, more specifically the *Arabidopsis thaliana* (At)CKX.

To effect expression of a protein in a cell, tissue or organ, preferably of plant origin, either the protein may be introduced directly to said cell, such as by microinjection or ballistic means or alternatively, an isolated nucleic acid molecule encoding said protein may be introduced into said cell, tissue or organ in an expressible format.

Preferably, the DNA sequence of the invention comprises a coding sequence or open reading frame (ORF) encoding a cytokinin oxidase protein or a homologue or derivative thereof or an immunologically active and/or functional fragment thereof as defined supra. The preferred protein of the invention comprises the amino acid sequence of said cytokinin oxidase. Preferably said cytokinin oxidase is a plant cytokinin oxidase and more specifically a *Arabidopsis thaliana* (At)CKX.

With "vector" or "vector sequence" is meant a DNA sequence which can be introduced in an organism by transformation and can be stably maintained in said organism. Vector maintenance is possible in e.g. cultures of *Escherichia coli*, *A. tumefaciens*,

Saccharomyces cerevisiae or Schizosaccharomyces pombe. Other vectors such as phagemids and cosmid vectors can be maintained and multiplied in bacteria and/or viruses. Vector sequences generally comprise a set of unique sites recognized by restriction enzymes, the multiple cloning site (MCS), wherein one or more non-vector sequence(s) can be inserted.

With "non-vector sequence" is accordingly meant a DNA sequence which is integrated in one or more of the sites of the MCS comprised within a vector.

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"Expression vectors" form a subset of vectors which, by virtue of comprising the appropriate regulatory or control sequences enable the creation of an expressible format for the inserted non-vector sequence(s), thus allowing expression of the protein encoded by said non-vector sequence(s). Expression vectors are known in the art enabling protein expression in organisms including bacteria (e.g. *E. coli*), fungi (e.g. *S. cerevisiae*, *S. pombe*, *Pichia pastoris*), insect cells (e.g. baculoviral expression vectors), animal cells (e.g. COS or CHO cells) and plant cells (e.g. potato virus X-based expression vectors).

The current invention clearly includes any cytokinin oxidase, homologue, derivative and/or immunologically active and/or functional fragment thereof as defined supra. Preferably said cytokinin oxidase is a plant cytokinin oxidase, more specifically a *Arabidopsis thaliana* (At)CKX.

As an alternative to expression vector-mediated protein production in biological systems, chemical protein synthesis can be applied. Synthetic peptides can be manufactured in solution phase or in solid phase. Solid phase peptide synthesis (Merrifield 1963) is, however, the most common way and involves the sequential addition of amino acids to create a linear peptide chain. Solid phase peptide synthesis includes cycles consisting of three steps: (i) immobilization of the carboxy-terminal amino acid of the growing peptide chain to a solid support or resin; (ii) chain assembly, a process consisting of activation, coupling and deprotection of the amino acid to be added to the growing peptide chain; and (iii) cleavage involving removal of the completed peptide chain from the resin and removal of the protecting groups from the amino acid side chains. Common approaches in solid phase peptide synthesis include Fmoc/tBu (9-fluorenylmethyloxycarbonyl/t-butyl) and Boc (t-butyloxycarbonyl) as the amino-terminal protecting groups of amino acids. Amino acid side chain protecting groups include methyl (Me), formyl (CHO), ethyl (Et), acetyl (Ac), t-butyl (t-Bu), anisyl, benzyl (Bzl), trifluroacetyl (Tfa), N-hydroxysuccinimide OSu), (ONSu, benzoyl (Bz), 4-methylbenzyl (Meb), thioanizyl, thiocresyl, benzyloxymethyl (Bom), 4-nitrophenyl (ONp), benzyloxycarbonyl (Z), 2-nitrobenzoyl

(NBz), 2-nitrophenylsulphenyl (Nps), 4-toluenesulphonyl (Tosyl,Tos), pentafluorophenyl (Pfp), diphenylmethyl (Dpm), 2-chlorobenzyloxycarbonyl (Cl-Z), 2,4,5-trichlorophenyl, 2-bromobenzyloxycarbonyl (Br-Z), tripheylmethyl (Trityl, Trt), and 2,5,7,8-pentamethyl-chroman-6-sulphonyl (Pmc). During chain assembly, Fmoc or Boc are removed resulting in an activated amino-terminus of the amino acid residue bound to the growing chain. The carboxy-terminus of the incoming amino acid is activated by conversion into a highly reactive ester, e.g. by HBTU. With current technologies (e.g. PerSeptive Biosystems 9050 synthesizer, Applied Biosystems Model 431A Peptide Synthesizer), linear peptides of up to 50 residues can be manufactured. A number of guidelines is available to produce peptides that are suitable for use in biological systems including (i) limiting the use of difficult amino acids such as cys, met, trp (easily oxidized and/or degraded during peptide synthesis) or arg; (ii) minimize hydrophobic amino acids (can impair peptide solubility); and (iii) prevent an amino-terminal glutamic acid (can cyclize to pyroglutamate).

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By "expressible format" is meant that the isolated nucleic acid molecule is in a form suitable for being transcribed into mRNA and/or translated to produce a protein, either constitutively or following induction by an intracellular or extracellular signal, such as an environmental stimulus or stress (mitogens, anoxia, hypoxia, temperature, sait, light, dehydration, etc) or a chemical compound such as IPTG (isopropyl-β-Dthiogalactopyranoside) or such as an antibiotic (tetracycline, ampicillin, rifampicin, kanamycin), hormone (e.g. gibberellin, auxin, cytokinin, glucocorticoid, brassinosteroid, ethylene, abscisic acid etc), hormone analogue (indolacetic acid (IAA), 2,4-D, etc), metal (zinc, copper, iron, etc), or dexamethasone, amongst others. As will be known to those skilled in the art, expression of a functional protein may also require one or more post-translational modifications, such glycosylation, as phosphorylation, dephosphorylation, or one or more protein-protein interactions, amongst others. All such processes are included within the scope of the term "expressible format".

Preferably, expression of a protein in a specific cell, tissue, or organ, preferably of plant origin, is effected by introducing and expressing an isolated nucleic acid molecule encoding said protein, such as a cDNA molecule, genomic gene, synthetic oligonucleotide molecule, mRNA molecule or open reading frame, to said cell, tissue or organ, wherein said nucleic acid molecule is placed operably in connection with suitable regulatory or control sequences including a promoter, preferably a plant-expressible promoter, and a terminator sequence.

Reference herein to a "promoter" is to be taken in its broadest context and includes the transcriptional regulatory sequences derived from a classical eukaryotic genomic gene, including the TATA box which is required for accurate transcription initiation, with or without a CCAAT box sequence and additional regulatory or control elements (i.e. upstream activating sequences, enhancers and silencers) which alter gene expression in response to developmental and/or external stimuli, or in a tissue-specific manner.

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The term "promoter" also includes the transcriptional regulatory sequences of a classical prokaryotic gene, in which case it may include a -35 box sequence and/or a -10 box transcriptional regulatory sequences.

The term "promoter" is also used to describe a synthetic or fusion molecule, or derivative which confers, activates or enhances expression of a nucleic acid molecule in a cell, tissue or organ.

Promoters may contain additional copies of one or more specific regulatory elements, to further enhance expression and/or to alter the spatial expression and/or temporal expression of a nucleic acid molecule to which it is operably connected. Such regulatory elements may be placed adjacent to a heterologous promoter sequence to drive expression of a nucleic acid molecule in response to e.g. copper, glucocorticoids, dexamethasone, tetracycline, gibberellin, cAMP, abscisic acid, auxin, wounding, ethylene, jasmonate or salicylic acid or to confer expression of a nucleic acid molecule to specific cells, tissues or organs such as meristems, leaves, roots, embryo, flowers, seeds or fruits.

In the context of the present invention, the promoter preferably is a plant-expressible promoter sequence. Promoters that also function or solely function in non-plant cells such as bacteria, yeast cells, insect cells and animal cells are not excluded from the invention. By "plant-expressible" is meant that the promoter sequence, including any additional regulatory elements added thereto or contained therein, is at least capable of inducing, conferring, activating or enhancing expression in a plant cell, tissue or organ, preferably a monocotyledonous or dicotyledonous plant cell, tissue, or organ.

Regulatable promoters as part of a binary viral plant expression system are also known to the skilled artisan (Yadav 1999 – WO9922003; Yadav 2000 – WO0017365).

The terms "plant-operable" and "operable in a plant" when used herein, in respect of a promoter sequence, shall be taken to be equivalent to a plant-expressible promoter sequence.

In the present context, a "regulatable promoter sequence" is a promoter that is capable of conferring expression on a structural gene in a particular cell, tissue, or organ or group of cells, tissues or organs of a plant, optionally under specific conditions, however does generally not confer expression throughout the plant under all conditions. Accordingly, a regulatable promoter sequence may be a promoter sequence that confers expression on a gene to which it is operably connected in a particular location within the plant or alternatively, throughout the plant under a specific set of conditions, such as following induction of gene expression by a chemical compound or other elicitor.

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Preferably, the regulatable promoter used in the performance of the present invention confers expression in a specific location within the plant, either constitutively or following induction, however not in the whole plant under any circumstances. Included within the scope of such promoters are cell-specific promoter sequences, tissue-specific promoter sequences, organ-specific promoter sequences, cell cycle specific gene promoter sequences, inducible promoter sequences and constitutive promoter sequences that have been modified to confer expression in a particular part of the plant at any one time, such as by integration of said constitutive promoter within a transposable genetic element (*Ac, Ds, Spm, En*, or other transposon).

The term "cell-specific" shall be taken to indicate that expression is predominantly in a particular cell or cell-type, preferably of plant origin, albeit not necessarily exclusively in said cell or cell-type.

Similarly, the term "tissue-specific" shall be taken to indicate that expression is predominantly in a particular tissue or tissue-type, preferably of plant origin, albeit not necessarily exclusively in said tissue or tissue-type.

Similarly, the term "organ-specific" shall be taken to indicate that expression is predominantly in a particular organ, preferably of plant origin, albeit not necessarily exclusively in said organ.

Similarly, the term "cell cycle specific" shall be taken to indicate that expression is predominantly cyclic and occurring in one or more, not necessarily consecutive phases of the cell cycle albeit not necessarily exclusively in cycling cells, preferably of plant origin.

Those skilled in the art will be aware that an "inducible promoter" is a promoter the transcriptional activity of which is increased or induced in response to a developmental, chemical, environmental, or physical stimulus. Similarly, the skilled craftsman will understand that a "constitutive promoter" is a promoter that is transcriptionally active

throughout most, but not necessarily all parts of an organism, preferably a plant, during most, but not necessarily all phases of its growth and development.

Those skilled in the art will readily be capable of selecting appropriate promoter sequences for use in regulating appropriate expression of the cytokinin oxidase protein from publicly-available or readily-available sources, without undue experimentation.

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Placing a nucleic acid molecule under the regulatory control of a promoter sequence, or in operable connection with a promoter sequence, means positioning said nucleic acid molecule such that expression is controlled by the promoter sequence. A promoter is usually, but not necessarily, positioned upstream, or at the 5'-end, and within 2 kb of the start site of transcription, of the nucleic acid molecule which it regulates. In the construction of heterologous promoter/structural gene combinations it is generally preferred to position the promoter at a distance from the gene transcription start site that is approximately the same as the distance between that promoter and the gene it controls in its natural setting (i.e., the gene from which the promoter is derived). As is known in the art, some variation in this distance can be accommodated without loss of promoter function. Similarly, the preferred positioning of a regulatory sequence element with respect to a heterologous gene to be placed under its control is defined by the positioning of the element in its natural setting (i.e., the gene from which it is derived). Again, as is known in the art, some variation in this distance can also occur.

Examples of promoters suitable for use in gene constructs of the present invention include those listed in Table 4, amongst others. The promoters listed in Table 4 are provided for the purposes of exemplification only and the present invention is not to be limited by the list provided therein. Those skilled in the art will readily be in a position to provide additional promoters that are useful in performing the present invention.

In the case of constitutive promoters or promoters that induce expression throughout the entire plant, it is preferred that such sequences are modified by the addition of nucleotide sequences derived from one or more of the tissue-specific promoters listed in Table 4, or alternatively, nucleotide sequences derived from one or more of the above-mentioned tissue-specific inducible promoters, to confer tissue-specificity thereon. For example, the CaMV 35S promoter may be modified by the addition of maize *Adh1* promoter sequence, to confer anaerobically-regulated root-specific expression thereon, as described previously (Ellis *et al.*, 1987). Another example describes conferring root specific or root abundant gene expression by fusing the CaMV35S promoter to elements

of the maize glycine-rich protein GRP3 gene (Feix and Wulff 2000 - WO0015662). Such modifications can be achieved by routine experimentation by those skilled in the art. The term "terminator" refers to a DNA sequence at the end of a transcriptional unit which signals termination of transcription. Terminators are 3'-non-translated DNA sequences containing a polyadenylation signal, which facilitates the addition of polyadenylate sequences to the 3'-end of a primary transcript. Terminators active in cells derived from viruses, yeasts, moulds, bacteria, insects, birds, mammals and plants are known and described in the literature. They may be isolated from bacteria, fungi, viruses, animals and/or plants.

Table 4. Exemplary plant-expressible promoters for use in the performance of the present invention

GENE SOURCE	EXPRESSION PATTERN	REFERENCE
α-amylase (<i>Amy32b</i>)	aleurone	Lanahan, M.B., et al., Plant Cell 4:203-211, 1992; Skriver, K., et al. Proc. Natl. Acad. Sci. (USA) 88: 7266-7270, 1991
cathepsin β-like gene	aleurone	Cejudo, F.J., et al. Plant Molecular Biology 20:849-856, 1992.
Agrobacterium rhizogenes rolB	cambium	Nilsson et al., Physiol. Plant. 100:456- 462, 1997
AtPRP4	flowers	http://salus.medium.edu/mmg/tierney/html
chalcone synthase (chsA)	flowers	Van der Meer, et al., Plant Mol. Biol. 15, 95-109, 1990.
LAT52	anther	Twell et al Mol. Gen Genet. 217:240- 245 (1989)
apetala-3	flowers	
chitinase	fruit (berries, grapes, etc)	Thomas et al. CSIRO Plant Industry, Urrbrae, South Australia, Australia; http://winetitles.com.au/gwrdc/csh95- 1.html
rbcs-3A	green tissue (eg leaf)	Lam, E. et al., The Plant Cell 2: 857-866, 1990.; Tucker et al., Plant Physiol. 113: 1303-1308, 1992.
leaf-specific genes	leaf	Baszczynski, et al., Nucl. Acid Res. 16: 4732, 1988.
AtPRP4	leaf	http://salus.medium.edu/mmg/tierney/
chlorella virus adenine methyltransferase gene promoter	leaf	Mitra and Higgins, 1994, Plant Molecular Biology 26: 85-93
aldP gene promoter from rice	leaf	Kagaya et al., 1995, Molecular and General Genetics 248: 668-674
rbcs promoter from rice or tomato	leaf	Kyozuka et al., 1993, Plant Physiology 102: 991-1000
Pinus cab-6	leaf	Yamamoto <i>et al., Plant Cell Physiol.</i> 35:773-778, 1994.
rubisco promoter	leaf	
cab (chlorophyll	leaf	

a/b/binding protein		
SAM22	senescent leaf	Crowell, et al., Plant Mol. Biol. 18:
		459-466, 1992.
Itp gene (lipid transfer gene)		Fleming, et al, Plant J. 2, 855-862.
R. japonicum nif gene	Nodule	United States Patent No. 4, 803, 165
B. japonicum nifH gene	Nodule	United States Patent No. 5, 008, 194
GmENOD40	Nodule	Yang, et al., The Plant J. 3: 573-585.
PEP carboxylase (PEPC)	Nodule	Pathirana, et al., Plant Mol. Biol. 20: 437-450, 1992.
leghaemoglobin (Lb)	Nodule	Gordon, et al., J. Exp. Bot. 44: 1453-1465, 1993.
Tungro bacilliform virus gene	phloem	Bhattacharyya-Pakrasi, et al, The Plant J. 4: 71-79, 1992.
pollen-specific genes	pollen; microspore	Albani, et al., Plant Mol. Biol. 15: 605, 1990; Albani, et al., Plant Mol. Biol. 16: 501, 1991)
Zm13	pollen	Guerrero et al Mol. Gen. Genet. 224:161-168 (1993)
apg gene	microspore	Twell et al Sex. Plant Reprod. 6:217-224 (1993)
maize pollen-specific gene	pollen	Hamilton, et al., Plant Mol. Biol. 18: 211-218, 1992.
sunflower pollen- expressed gene	pollen	Baltz, et al., The Plant J. 2: 713-721, 1992.
B. napus pollen- specific gene	pollen;anther; tapetum	Arnoldo, et al., J. Cell. Biochem., Abstract No. Y101, 204, 1992.
root-expressible genes	roots	Tingey, et al., EMBO J. 6: 1, 1987.
tobacco auxin-inducible gene	root tip	Van der Zaal, et al., Plant Mol. Biol. 16, 983, 1991.
β-tubulin	root	Oppenheimer, et al., Gene 63: 87, 1988.
tobacco root-specific genes	root	Conkling, et al., Plant Physiol. 93: 1203, 1990.
B. napus G1-3b gene	root	United States Patent No. 5, 401, 836
SbPRP1	roots	Suzuki et al., Plant Mol. Biol. 21: 109- 119, 1993.
AtPRP1; AtPRP3	roots; root hairs	http://salus.medium.edu/mmg/tierney/ html
RD2 gene	root cortex	http://www2.cnsu.edu/ncsu/research
TobRB7 gene	root vasculature	http://www2.cnsu.edu/ncsu/research
AtPRP4	leaves; flowers; lateral root primordia	http://salus.medium.edu/mmg/tierney/ html
seed-specific genes	seed	Simon, et al., Plant Mol. Biol. 5: 191,

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		1985; Scofield, et al., J. Biol. Chem. 262: 12202, 1987.; Baszczynski, et al., Plant Mol. Biol. 14: 633, 1990.
Brazil Nut albumin	seed	Pearson, et al., Plant Mol. Biol. 18: 235-245, 1992.
legumin	seed	Ellis, et al., Plant Mol. Biol. 10: 203- 214, 1988.
glutelin (rice)	seed	Takaiwa, et al., Mol. Gen. Genet. 208: 15-22, 1986; Takaiwa, et al., FEBS Letts. 221: 43-47, 1987.
zein	seed	Matzke et al Plant Mol Biol, 14(3):323-32 1990
napA	seed	Stalberg, et al, Planta 199: 515-519, 1996.
wheat LMW and HMW glutenin-1	endosperm	Mol Gen Genet 216:81-90, 1989; NAR 17:461-2, 1989
wheat SPA	seed	Albani <i>et al</i> , Plant Cell, 9: 171-184, 1997
wheat α , β , γ -gliadins	endosperm	EMBO 3:1409-15, 1984
barley Itr1 promoter	endosperm	
barley B1, C, D, hordein	endosperm	Theor Appl Gen 98:1253-62, 1999; Plant J 4:343-55, 1993; Mol Gen Genet 250:750-60, 1996
barley DOF	endosperm	Mena et al, The Plant Journal, 116(1): 53-62, 1998
blz2	endosperm	EP99106056.7
synthetic promoter	endosperm	Vicente-Carbajosa et al., Plant J. 13: 629-640, 1998.
rice prolamin NRP33	endosperm	Wu et al, Plant Cell Physiology 39(8) 885-889, 1998
rice α-globulin Glb-1	endosperm	Wu et al, Plant Cell Physiology 39(8) 885-889, 1998
rice OSH1	embryo	Sato et al, Proc. Natl. Acad. Sci. USA, 93: 8117-8122, 1996
rice α-globulin REB/OHP-1	endosperm	Nakase et al. Plant Mol. Biol. 33: 513-522, 1997
rice ADP-glucose PP	endosperm	Trans Res 6:157-68, 1997
maize ESR gene family	endosperm	Plant J 12:235-46, 1997
sorgum γ-kafirin	endosperm	PMB 32:1029-35, 1996
KNOX	embryo	Postma-Haarsma et al, Plant Mol. Biol. 39:257-71, 1999
rice oleosin	embryo and aleuron	Wu et at, J. Biochem., 123:386, 1998
sunflower oleosin	seed (embryo and dry seed)	Cummins, et al., Plant Mol. Biol. 19: 873-876, 1992
LEAFY	shoot meristem	Weigel et al., Cell 69:843-859, 1992.

Arabidopsis thaliana knat1	shoot meristem	Accession number AJ131822
Malus domestica kn1	shoot meristem	Accession number Z71981
CLAVATA1	shoot meristem	Accession number AF049870
stigma-specific genes	stigma	Nasrallah, et al., Proc. Natl. Acad. Sci. USA 85: 5551, 1988; Trick, et al., Plant Mol. Biol. 15: 203, 1990.
class I patatin gene	tuber	Liu et al., Plant Mol. Biol. 153:386-395, 1991.
PCNA rice	meristem	Kosugi et al, Nucleic Acids Research 19:1571-1576, 1991; Kosugi S. and Ohashi Y, Plant Cell 9:1607-1619, 1997.
Pea TubA1 tubulin	Dividing cells	Stotz and Long, <i>Plant Mol.Biol.</i> 41, 601-614. 1999
Arabidopsis cdc2a	cycling cells	Chung and Parish, FEBS Lett, 3;362(2):215-9, 1995
Arabidopsis Rop1A	Anthers; mature pollen + pollen tubes	Li et al. 1998 <i>Plant Physiol</i> 118, 407-417.
Arabidopsis AtDMC1	Meiosis-associated	Klimyuk and Jones 1997 Plant J. 11, 1-14.
Pea PS-IAA4/5 and PS-IAA6	Auxin-inducible	Wong et al. 1996 <i>Plant J.</i> 9, 587-599.
Pea farnesyltransferase	Meristematic tissues; phloem near growing tissues; light- and sugar-repressed	Zhou et al. 1997 <i>Plant J.</i> 12, 921-930
Tobacco (<i>N. sylvestris</i>) cyclin B1;1	Dividing cells / meristematic tissue	Trehin et al. 1997 <i>Plant Mol.Biol.</i> 35, 667-672.
Catharanthus roseus Mitotic cyclins CYS (A- type) and CYM (B-type)	Dividing cells / meristematic tissue	Ito et al. 1997 <i>Plant J</i> . 11, 983-992
Arabidopsis cyc1At (=cyc B1;1) and cyc3aAt (A-type)	Dividing cells / meristematic tissue	Shaul et al. 1996 <i>Proc.Natl.Acad.Sci.U.S.A</i> 93, 4868- 4872.
Arabidopsis tef1 promoter box	Dividing cells / meristematic tissue	Regad et al. 1995 <i>Mol.Gen.Genet.</i> 248, 703-711.
Catharanthus roseus cyc07	Dividing cells / meristematic tissue	Ito et al. 1994 <i>Plant Mol.Biol.</i> 24, 863-878.

Table 4 (continued). Exemplary plant-expressible promoters for use in the performance of the present invention

11: 1	II: EXEMPLARY CONSTITUTIVE PROMOTERS	
GENE SOURCE	EXPRESSION PATTERN	REFERENCE
Actin	constitutive	McElroy et al, Plant Cell, 2: 163- 171, 1990
CAMV 35S	constitutive	Odell <i>et al,</i> Nature, 313: 810-812, 1985
CaMV 19S	constitutive	Nilsson <i>et al.</i> , Physiol. Plant. 100:456-462, 1997
GOS2	constitutive	de Pater <i>et al</i> , Plant J. 2:837-844, 1992
ubiquitin	constitutive	Christensen <i>et al,</i> Plant Mol. Biol. 18: 675-689, 1992
rice cyclophilin	constitutive	Buchholz <i>et al,</i> Plant Mol Biol. 25: 837-843, 1994
maize histone H3	constitutive	Lepetit <i>et al,</i> Mol. Gen. Genet. 231:276-285, 1992
alfalfa histone H3	constitutive	Wu <i>et al.</i> , Nucleic Acids Res. 17: 3057-3063, 1989; Wu <i>et al.</i> , Plant Mol. Biol. 11:641-649, 1988
actin 2	constitutive	An <i>et al</i> , Plant J. 10(1); 107-121, 1996

Table 4 (continued). Exemplary plant-expressible promoters for use in the performance of the present invention

III: EXEMPLARY STRESS-INDUCIBLE PROMOTERS		
NAME	STRESS	REFERENCE
P5CS (delta(1)- pyrroline-5-carboxylate syntase)	salt, water	Zhang et al. Plant Science. 129: 81- 89, 1997
cor15a	cold	Hajela et al., Plant Physiol. 93: 1246- 1252, 1990
cor15b	cold	Wlihelm et al., Plant Mol Biol. 23:1073-1077, 1993
cor15a (-305 to +78 nt)	cold, drought	Baker et al., Plant Mol Biol. 24: 701-713, 1994
rd29	salt, drought, cold	Kasuga et al., Nature Biotechnology 18:287-291, 1999
heat shock proteins, including artificial promoters containing the heat shock element (HSE)	heat	Barros et al., Plant Mol Biol 19: 665- 75, 1992. Marrs et al., Dev Genet.14: 27-41, 1993. Schoffl et al., Mol Gen Gent, 217: 246-53, 1989.
smHSP (small heat shock proteins)	heat	Waters et al, J Experimental Botany 47:325-338, 1996
wcs120	cold	Ouellet et al., FEBS Lett. 423: 324- 328, 1998
ci7	cold	Kirch et al., Plant Mol Biol 33: 897- 909, 1997
Adh	cold, drought, hypoxia	Dolferus et al., Plant Physiol 105: 1075-87, 1994
pwsi18	water: salt and drought	Joshee et al., Plant Cell Physiol 39: 64-72, 1998
ci21A	cold	Schneider et al., Plant Physiol 113: 335-45, 1997
Trg-31	drought	Chaudhary et al., Plant Mol Biol 30: 1247-57, 1996
osmotin	osmotic	Raghothama et al., Plant Mol Biol 23: 1117-28, 1993
Rab17	osmotic, ABA	Vilardell et al., Plant Mol Biol 17: 985-93, 1991
lapA	wounding, environmental	WO99/03977 University of California/INRA

Table 4 (continued). Exemplary plant-expressible promoters for use in the performance of the present invention

IV: EXEMPLARY PATHOGEN-INDUCIBLE PROMOTERS		
NAME	PATHOGEN	REFERENCE
RB7	Root-knot nematodes (Meloidogyne spp.)	US5760386 - North Carolina State University; Opperman et al (1994) Science 263: 221-23.
PR-1, 2, 3, 4, 5, 8, 11	fungal, viral, bacterial	Ward et al (1991) Plant Cell 3: 1085-1094; Reiss et al 1996; Lebel et al (1998), Plant J, 16(2):223-33; Melchers et al (1994), Plant J, 5(4):469-80; Lawton et al (1992), Plant Mol Biol, 19(5):735-43.
HMG2	nematodes	WO9503690 - Virginia Tech Intellectual Properties Inc .
Abi3	Cyst nematodes (Heterodera spp.)	Unpublished
ARM1	nematodes	Barthels et al., (1997) The Plant Cell 9, 2119-2134.
		WO 98/31822 - Plant Genetic Systems
Att0728	nematodes	Barthels et al., (1997) The Plant Cell 9, 2119-2134.
		PCT/EP98/07761
Att1712	nematodes	Barthels et al., (1997) The Plant Cell 9, 2119-2134.
		PCT/EP98/07761
Gst1	Different types of pathogens	Strittmatter et al (1996) Mol. Plant-Microbe Interact. 9, 68-73.
LEMMI	nematodes	WO 92/21757 – Plant Genetic Systems
CLE	geminivirus	PCT/EP99/03445 - CINESTAV
PDF1.2	Fungal including Alternaria brassicicola and Botrytis cinerea	Manners et al (1998), Plant Mol Biol, 38(6):1071-80.
Thi2.1	Fungal – Fusarium oxysporum f sp. matthiolae	Vignutelli et al (1998) Plant J;14(3):285-95
DB#226	nematodes	Bird and Wilson (1994) Mol. Plant- Microbe Interact., 7, 419-42 WO 95.322888

DB#280	nematodes	Bird and Wilson (1994) Mol. Plant- Microbe Interact., 7, 419-42 WO 95.322888
Cat2	nematodes	Niebel et al (1995) Mol Plant Microbe Interact 1995 May- Jun;8(3):371-8
□Tub	nematodes	Aristizabal et al (1996), 8 th International Congress on Plant- Microbe Interaction, Knoxville US B-29
SHSP	nematodes	Fenoll et al (1997) In: Cellular and molecular aspects of plant- nematode interactions. Kluwer Academic, C. Fenoll, F.M.W. Grundler and S.A. Ohl (Eds.),
Tsw12	nematodes	Fenoli et al (1997) In: Cellular and molecular aspects of plant- nematode interactions. Kluwer Academic, C. Fenoll, F.M.W. Grundler and S.A. Ohl (Eds.)
Hs1(pro1)	nematodes	WO 98/122335 - Jung
NsLTP	viral, fungal, bacterial	Molina & Garc´ia-Olmedo (1993) FEBS Lett, 316(2):119-22
RIP	viral, fungal	Tumer et al (1997) Proc Natl Acad Sci U S A, 94(8):3866-71

Examples of terminators particularly suitable for use in the gene constructs of the present invention include the *Agrobacterium tumefaciens* nopaline synthase (NOS) gene terminator, the *Agrobacterium tumefaciens* octopine synthase (OCS) gene terminator sequence, the Cauliflower mosaic virus (CaMV) 35S gene terminator sequence, the *Oryza sativa* ADP-glucose pyrophosphorylase terminator sequence (t3'Bt2), the *Zea mays* zein gene terminator sequence, the *rbcs-1A* gene terminator, and the *rbcs-3A* gene terminator sequences, amongst others.

Preferred promoter sequences of the invention include root specific promoters such as but not limited to the ones listed in Table 5 and as outlined in the Examples.

Table 5. Exemplary root specific promoters for use in the performance of the present invention

NAME	ORIGIN	REFERENCE
SbPRP1	Soybean	Suzuki et al., Plant Mol Biol, 21: 109-119, 1993
636 bp fragment of TobRB7	Tobacco	Yamamoto et al., Plant Cell 3:371-382, 1991
GGPS3	Arabidopsis	Okada et al., Plant Physiol 122: 1045-1056, 2000
580 bp fragment of prxEa	Arabidopsis	Wanapu and Shinmyo, Ann N Y Acad Sci 782: 107-114, 1996
lds2 promoter	Barley	Okumura et al., Plant Mol Biol 25: 705-719, 1994
AtPRP3	Arabidopsis	Fowler et al., Plant Physiol 121: 1081-1092, 1999

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Those skilled in the art will be aware of additional promoter sequences and terminator sequences which may be suitable for use in performing the invention. Such sequences may readily be used without any undue experimentation.

In the context of the current invention, "ectopic expression" or "ectopic overexpression" of a gene or a protein are conferring to expression patterns and/or expression levels of said gene or protein normally not occurring under natural conditions, more specifically is meant increased expression and/or increased expression levels. Ectopic expression can be achieved in a number of ways including operably linking of a coding sequence encoding said protein to an isolated homologous or heterologous promoter in order to create a chimeric gene and/or operably linking said coding sequence to its own isolated promoter (i.e. the unisolated promoter naturally driving expression of said protein) in order to create a recombinant gene duplication or gene multiplication effect. With "ectopic co-expression" is meant the ectopic expression or ectopic overexpression of two or more genes or proteins. The same or, more preferably, different promoters are used to confer ectopic expression of said genes or proteins.

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Preferably, the promoter sequence used in the context of the present invention is operably linked to a coding sequence or open reading frame (ORF) encoding a cytokinin

oxidase protein or a homologue, derivative or an immunologically active and/or functional fragment thereof as defined supra.

"Downregulation of expression" as used herein means lowering levels of gene expression and/or levels of active gene product and/or levels of gene product activity. Decreases in expression may be accomplished by e.g. the addition of coding sequences or parts thereof in a sense orientation (if resulting in co-suppression) or in an antisense orientation relative to a promoter sequence and furthermore by e.g. insertion mutagenesis (e.g. T-DNA insertion or transposon insertion) or by gene silencing strategies as described by e.g. Angell and Baulcombe (1998 - WO9836083), Lowe et al. (1989 - WO9853083), Lederer et al. (1999 - WO9915682) or Wang et al. (1999 - WO9953050). Genetic constructs aimed at silencing gene expression may have the nucleotide sequence of said gene (or one or more parts thereof) contained therein in a sense and/or antisense orientation relative to the promoter sequence. Another method to downregulate gene expression comprises the use of ribozymes.

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Modulating, including lowering, the level of active gene products or of gene product activity can be achieved by administering or exposing cells, tissues, organs or organisms to said gene product, a homologue, derivative and/or immunologically active fragment thereof. Immunomodulation is another example of a technique capable of downregulation levels of active gene product and/or of gene product activity and comprises administration of or exposing to or expressing antibodies to said gene product to or in cells, tissues, organs or organisms wherein levels of said gene product and/or gene product activity are to be modulated. Such antibodies comprise "plantibodies", single chain antibodies, IgG antibodies and heavy chain camel antibodies as well as fragments thereof.

Modulating, including lowering, the level of active gene products or of gene product activity can futhermore be achieved by administering or exposing cells, tissues, organs or organisms to an agonist of said gene product or the activity thereof. Such agonists include proteins (comprising e.g. kinases and proteinases) and chemical compounds identified according to the current invention as described supra.

In the context of the current invention is envisaged the downregulation of the expression of a cytokinin oxidase gene as defined higher. Preferably said cytokinin oxidase gene is a plant cytokinin oxidase gene, more specifically an *AtCKX*. The invention further comprises downregulation of levels of a cytokinin oxidase protein or of a cytokinin oxidase activity whereby said cytokinin oxidase protein has been defined supra.

Preferably said cytokinin oxidase protein is a plant cytokinin oxidase, more specifically an AtCKX.

By "modifying cell fate and/or plant development and/or plant morphology and/or biochemistry and/or physiology" is meant that one or more developmental and/or morphological and/or biochemical and/or physiological characteristics of a plant is altered by the performance of one or more steps pertaining to the invention described herein.

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"Cell fate" refers to the cell-type or cellular characteristics of a particular cell that are produced during plant development or a cellular process therefor, in particular during the cell cycle or as a consequence of a cell cycle process.

"Plant development" or the term "plant developmental characteristic" or similar term shall, when used herein, be taken to mean any cellular process of a plant that is involved in determining the developmental fate of a plant cell, in particular the specific tissue or organ type into which a progenitor cell will develop. Cellular processes relevant to plant development will be known to those skilled in the art. Such processes include, for example, morphogenesis, photomorphogenesis, shoot development, root development, vegetative development, reproductive development, stem elongation, flowering, and regulatory mechanisms involved in determining cell fate, in particular a process or regulatory process involving the cell cycle.

"Plant morphology" or the term "plant morphological characteristic" or similar term will, when used herein, be understood by those skilled in the art to refer to the external appearance of a plant, including any one or more structural features or combination of structural features thereof. Such structural features include the shape, size, number, position, colour, texture, arrangement, and patternation of any cell, tissue or organ or groups of cells, tissues or organs of a plant, including the root, stem, leaf, shoot, petiole, trichome, flower, petal, stigma, style, stamen, pollen, ovule, seed, embryo, endosperm, seed coat, aleurone, fibre, fruit, cambium, wood, heartwood, parenchyma, aerenchyma, sieve element, phloem or vascular tissue, amongst others.

"Plant biochemistry" or the term "plant biochemical characteristic" or similar term will, when used herein, be understood by those skilled in the art to refer to the metabolic and catalytic processes of a plant, including primary and secondary metabolism and the products thereof, including any small molecules, macromolecules or chemical compounds, such as but not limited to starches, sugars, proteins, peptides, enzymes, hormones, growth factors, nucleic acid molecules, celluloses, hemicelluloses, calloses,

lectins, fibres, pigments such as anthocyanins, vitamins, minerals, micronutrients, or macronutrients, that are produced by plants.

"Plant physiology" or the term "plant physiological characteristic" or similar term will, when used herein, be understood to refer to the functional processes of a plant, including developmental processes such as growth, expansion and differentiation, sexual development, sexual reproduction, seed set, seed development, grain filling, reproduction, cell division, dormancy, germination, light adaptation, asexual photosynthesis, leaf expansion, fibre production, secondary growth or wood production, amongst others; responses of a plant to externally-applied factors such as metals, chemicals, hormones, growth factors, environment and environmental stress factors (eg. anoxia, hypoxia, high temperature, low temperature, dehydration, light, daylength, flooding, salt, heavy metals, amongst others), including adaptive responses of plants to said externally-applied factors.

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Means for introducing recombinant DNA into plant tissue or cells include, but are not limited to, transformation using CaCl2 and variations thereof, in particular the method 15 described by Hanahan (1983), direct DNA uptake into protoplasts (Krens et al, 1982; Paszkowski et al, 1984), PEG-mediated uptake to protoplasts (Armstrong et al, 1990) microparticle bombardment, electroporation (Fromm et al., 1985), microinjection of DNA (Crossway et al., 1986), microparticle bombardment of tissue explants or cells (Christou et al, 1988; Sanford, 1988), vacuum-infiltration of tissue with nucleic acid, or in the case 20 of plants, T-DNA-mediated transfer from Agrobacterium to the plant tissue as described essentially by An et al.(1985), Dodds et al., (1985), Herrera-Estrella et al. (1983a, 1983b, 1985). Methods for transformation of monocotyledonous plants are well known in the art and include Agrobacterium-mediated transformation (Cheng et al., 1997 - WO9748814; Hansen 1998 - WO9854961; Hiei et al., 1994 - WO9400977; Hiei et al., 1998 -25 WO9817813; Rikiishi et al., 1999 - WO9904618; Saito et al., 1995 - WO9506722), microprojectile bombardment (Adams et al., 1999 - US5969213; Bowen et al., 1998 -US5736369; Chang et al., 1994 - WO9413822; Lundquist et al., 1999 -US5874265/US5990390; Vasil and Vasil, 1995 - US5405765. Walker et al., 1999 -US5955362), DNA uptake (Eyal et al., 1993 - WO9318168), microinjection of Agrobacterium cells (von Holt, 1994 - DE4309203) and sonication (Finer et al., 1997 -US5693512).

For microparticle bombardment of cells, a microparticle is propelled into a cell to produce a transformed cell. Any suitable ballistic cell transformation methodology and apparatus can be used in performing the present invention. Exemplary apparatus and procedures are disclosed by Stomp *et al.* (U.S. Patent No. 5,122,466) and Sanford and Wolf (U.S. Patent No. 4,945,050). When using ballistic transformation procedures, the gene construct may incorporate a plasmid capable of replicating in the cell to be transformed. Examples of microparticles suitable for use in such systems include 1 to 5 µm gold spheres. The DNA construct may be deposited on the microparticle by any suitable technique, such as by precipitation.

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A whole plant may be regenerated from the transformed or transfected cell, in accordance with procedures well known in the art. Plant tissue capable of subsequent clonal propagation, whether by organogenesis or embryogenesis, may be transformed with a gene construct of the present invention and a whole plant regenerated therefrom. The particular tissue chosen will vary depending on the clonal propagation systems available for, and best suited to, the particular species being transformed. Exemplary tissue targets include leaf disks, pollen, embryos, cotyledons, hypocotyls, megagametophytes, callus tissue, existing meristematic tissue (e.g., apical meristem, axillary buds, and root meristems), and induced meristem tissue (e.g., cotyledon meristem and hypocotyl meristem).

The term "organogenesis", as used herein, means a process by which shoots and roots are developed sequentially from meristematic centres.

The term "embryogenesis", as used herein, means a process by which shoots and roots develop together in a concerted fashion (not sequentially), whether from somatic cells or gametes.

Preferably, the plant is produced according to the inventive method is transfected or transformed with a genetic sequence, or amenable to the introduction of a protein, by any art-recognized means, such as microprojectile bombardment, microinjection, *Agrobacterium*-mediated transformation (including *in planta* transformation), protoplast fusion, or electroporation, amongst others. Most preferably said plant is produced by *Agrobacterium*-mediated transformation.

Agrobacterium-mediated transformation or agrolistic transformation of plants, yeast, moulds or filamentous fungi is based on the transfer of part of the transformation vector sequences, called the T-DNA, to the nucleus and on integration of said T-DNA in the genome of said eukaryote.

With "Agrobacterium" is meant a member of the Agrobacteriaceae, more preferably Agrobacterium or Rhizobacterium and most preferably Agrobacterium tumefaciens.

With "T-DNA", or transferred DNA, is meant that part of the transformation vector flanked by T-DNA borders which is, after activation of the *Agrobacterium vir* genes, nicked at the T-DNA borders and is transferred as a single stranded DNA to the nucleus of an eukaryotic cell.

When used herein, with "T-DNA borders", "T-DNA border region", or "border region" are meant either right T-DNA border (RB) or left T-DNA border (LB). Such a border comprises a core sequence flanked by a border inner region as part of the T-DNA flanking the border and/or a border outer region as part of the vector backbone flanking the border. The core sequences comprise 22 bp in case of octopine-type vectors and 25 bp in case of nopaline-type vectors. The core sequences in the right border region and left border region form imperfect repeats. Border core sequences are indispensable for recognition and processing by the *Agrobacterium* nicking complex consisting of at least VirD1 and VirD2. Core sequences flanking a T-DNA are sufficient to promote transfer of said T-DNA. However, efficiency of transformation using transformation vectors carrying said T-DNA solely flanked by said core sequences is low. Border inner and outer regions are known to modulate efficiency of T-DNA transfer (Wang et al. 1987). One element enhancing T-DNA transfer has been characterized and resides in the right border outer region and is called *overdrive* (Peralta et al. 1986, van Haaren et al. 1987).

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With "T-DNA transformation vector" or "T-DNA vector" is meant any vector encompassing a T-DNA sequence flanked by a right and left T-DNA border consisting of at least the right and left border core sequences, respectively, and used for transformation of any eukaryotic cell.

With "T-DNA vector backbone sequence" or "T-DNA vector backbone sequences" is meant all DNA of a T-DNA containing vector that lies outside of the T-DNA borders and, more specifically, outside the nicking sites of the border core imperfect repeats.

The current invention includes optimized T-DNA vectors such that vector backbone integration in the genome of a eukaryotic cell is minimized or absent. With "optimized T-DNA vector" is meant a T-DNA vector designed either to decrease or abolish transfer of vector backbone sequences to the genome of a eukaryotic cell. Such T-DNA vectors are known to the one familiar with the art and include those described by Hanson et al. (1999) and by Stuiver et al. (1999 - WO9901563).

The current invention clearly considers the inclusion of a DNA sequence encoding a cytokinin oxidase, homologue, derivative or immunologically active and/or functional fragment thereof as defined supra, in any T-DNA vector comprising binary

transformation vectors, super-binary transformation vectors, co-integrate transformation vectors, Ri-derived transformation vectors as well as in T-DNA carrying vectors used in agrolistic transformation. Preferably, said cytokinin oxidase is a plant cytokinin oxidase, more specifically an *Arabidopsis thaliana* (At)CKX.

With "binary transformation vector" is meant a T-DNA transformation vector comprising:

- (a) a T-DNA region comprising at least one gene of interest and/or at least one selectable marker active in the eukaryotic cell to be transformed; and
- (b) a vector backbone region comprising at least origins of replication active in *E. coli* and *Agrobacterium* and markers for selection in *E. coli* and *Agrobacterium*.
- The T-DNA borders of a binary transformation vector can be derived from octopine-type or nopaline-type Ti plasmids or from both. The T-DNA of a binary vector is only transferred to a eukaryotic cell in conjunction with a helper plasmid.

With "helper plasmid" is meant a plasmid that is stably maintained in *Agrobacterium* and is at least carrying the set of *vir* genes necessary for enabling transfer of the T-DNA. Said set of *vir* genes can be derived from either octopine-type or nopaline-type Ti

plasmids or from both.

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With "super-binary transformation vector" is meant a binary transformation vector additionally carrying in the vector backbone region a *vir* region of the Ti plasmid pTiBo542 of the super-virulent *A. tumefaciens* strain A281 (EP0604662, EP0687730).

Super-binary transformation vectors are used in conjunction with a helper plasmid.

With "co-integrate transformation vector" is meant a T-DNA vector at least comprising:

- (a) a T-DNA region comprising at least one gene of interest and/or at least one selectable marker active in plants; and
- (b) a vector backbone region comprising at least origins of replication active in Escherichia coli and Agrobacterium, and markers for selection in E. coli and Agrobacterium, and a set of vir genes necessary for enabling transfer of the T-DNA.

The T-DNA borders and said set of *vir* genes of a said T-DNA vector can be derived from either octopine-type or nopaline-type Ti plasmids or from both.

With "Ri-derived plant transformation vector" is meant a binary transformation vector in which the T-DNA borders are derived from a Ti plasmid and said binary transformation vector being used in conjunction with a 'helper' Ri-plasmid carrying the necessary set of vir genes.

As used herein, the term "selectable marker gene" or "selectable marker" or "marker for selection" includes any gene which confers a phenotype on a cell in which it is expressed to facilitate the identification and/or selection of cells which are transfected or transformed with a gene construct of the invention or a derivative thereof. Suitable selectable marker genes contemplated herein include the ampicillin resistance (Amp^r), tetracycline resistance gene (Tc^r), bacterial kanamycin resistance gene (Kan^r), phosphinothricin resistance gene, neomycin phosphotransferase gene (*npt*II), hygromycin resistance gene, β-glucuronidase (GUS) gene, chloramphenicol acetyltransferase (CAT) gene, green fluorescent protein (*gfp*) gene (Haseloff *et al*, 1997), and luciferase gene, amongst others.

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With "agrolistics", "agrolistic transformation" or "agrolistic transfer" is meant here a transformation method combining features of *Agrobacterium*-mediated transformation and of biolistic DNA delivery. As such, a T-DNA containing target plasmid is co-delivered with DNA/RNA enabling in planta production of VirD1 and VirD2 with or without VirE2 (Hansen and Chilton 1996; Hansen et al. 1997; Hansen and Chilton 1997 - WO9712046).

With "foreign DNA" is meant any DNA sequence that is introduced in the host's genome by recombinant techniques. Said foreign DNA includes e.g. a T-DNA sequence or a part thereof such as the T-DNA sequence comprising the selectable marker in an expressible format. Foreign DNA furthermore include intervening DNA sequences as defined supra.

With "recombination event" is meant either a site-specific recombination event or a recombination event effected by transposon 'jumping'.

With "recombinase" is meant either a site-specific recombinase or a transposase.

With "recombination site" is meant either site-specific recombination sites or transposon border sequences.

With "site specific recombination event" is meant an event catalyzed by a system generally consisting of three elements: a pair of DNA sequences (the site-specific recombination sequences or sites) and a specific enzyme (the site-specific recombinase). The site-specific recombinase catalyzes a recombination reaction only between two site-specific recombination sequences depending on the orientation of the site-specific recombination sequences. Sequences intervening between two site-specific recombination sites will be inverted in the presence of the site-specific recombinase when the site-specific recombination sequences are oriented in opposite directions relative to one another (i.e. inverted repeats). If the site-specific recombination

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sequences are oriented in the same direction relative to one another (i.e. direct repeats), then any intervening sequences will be deleted upon interaction with the site-specific recombinase. Thus, if the site-specific recombination sequences are present as direct repeats at both ends of a foreign DNA sequence integrated into a eukaryotic genome, such integration of said sequences can subsequently be reversed by interaction of the site-specific recombination sequences with the corresponding site specific recombinase. A number of different site specific recombinase systems can be used including but not limited to the Cre/lox system of bacteriophage P1, the FLP/FRT system of yeast, the Gin recombinase of phage Mu, the Pin recombinase of E. coli, the PinB, PinD and PinF from Shigella, and the R/RS system of the pSR1 plasmid. Recombinases generally are integrases, resolvases or flippases. Also dual-specific recombinases can be used in conjunction with direct or indirect repeats of two different site-specific recombination sites corresponding to the dual-specific recombinase (WO99/25840). The two preferred site-specific recombinase systems are the bacteriophage P1 Cre/lox and the yeast FLP/FRT systems. In these systems a recombinase (Cre or FLP) interact specifically with its respective site-specific recombination sequence (lox or FRT respectively) to invert or excise the intervening sequences. The site-specific recombination sequences for each of these two systems are relatively short (34 bp for lox and 47 bp for FRT). Some of these systems have already been used with high efficiency in plants such as tobacco (Dale et al. 1990) and Arabidopsis (Osborne et al. 1995). Site-specific recombination systems have many applications in plant molecular biology including methods for control of homologous recombination (e.g. US5527695), for targeted insertion, gene stacking, etc. (WO99/25821) and for resolution of complex T-DNA integration patterns or for excision of a selectable marker (WO99/23202).

Although the site-specific recombination sequences must be linked to the ends of the DNA to be excised or to be inverted, the gene encoding the site specific recombinase may be located elsewhere. For example, the recombinase gene could already be present in the eukaryote's DNA or could be supplied by a later introduced DNA fragment either introduced directly into cells, through crossing or through cross-pollination. Alternatively, a substantially purified recombinase protein could be introduced directly into the eukaryotic cell, e.g. by micro-injection or particle bombardment. Typically, the site-specific recombinase coding region will be operably linked to regulatory sequences enabling expression of the site-specific recombinase in the eukaryotic cell.

With "recombination event effected by transposon jumping" or "transposase-mediated recombination" is meant a recombination event catalyzed by a system consisting of three elements: a pair of DNA sequences (the transposon border sequences) and a specific enzyme (the transposase). The transposase catalyzes a recombination reaction only between two transposon border sequences which are arranged as inverted repeats. A number of different transposon/transposase systems can be used including but not limited to the Ds/Ac system, the Spm system and the Mu system. These systems originate from corn but it has been shown that at least the Ds/Ac and the Spm system also function in other plants (Fedoroff et al. 1993, Schlappi et al. 1993, Van Sluys et al. 1987). Preferred are the Ds- and the Spm-type transposons which are delineated by 11 bp- and 13 bp- border sequences, respectively.

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Although the transposon border sequences must be linked to the ends of the DNA to be excised, the gene encoding the transposase may be located elsewhere. For example, the recombinase gene could already be present in the eukaryote's DNA or could be supplied by a later introduced DNA fragment either introduced directly into cells, through crossing or through cross-pollination. Alternatively, a substantially purified transposase protein could be introduced directly into cells, e.g. by microinjection or by particle bombardment.

As part of the current invention, transposon border sequences are included in a foreign DNA sequence such that they lie outside said DNA sequence and transform said DNA into a transposon-like entity that can move by the action of a transposase.

As transposons often reintegrate at another locus of the host's genome, segregation of the progeny of the hosts in which the transposase was allowed to act might be necessary to separate transformed hosts containing e.g. only the transposon footprint and transformed hosts still containing the foreign DNA.

In performing the present invention, the genetic element is preferably induced to mobilize, such as, for example, by the expression of a recombinase protein in the cell which contacts the integration site of the genetic element and facilitates a recombination event therein, excising the genetic element completely, or alternatively, leaving a "footprint", generally of about 20 nucleotides in length or greater, at the original integration site. Those hosts and host parts that have been produced according to the inventive method can be identified by standard nucleic acid hybridization and/or amplification techniques to detect the presence of the mobilizable genetic element or a gene construct comprising the same. Alternatively, in the case of transformed host cells,

tissues, and hosts wherein the mobilizable genetic element has been excised, it is possible to detect a footprint in the genome of the host which has been left following the excision event, using such techniques. As used herein, the term "footprint" shall be taken to refer to any derivative of a mobilizable genetic element or gene construct comprising the same as described herein which is produced by excision, deletion or other removal of the mobilizable genetic element from the genome of a cell transformed previously with said gene construct. A footprint generally comprises at least a single copy of the recombination loci or transposon used to promote excision. However, a footprint may comprise additional sequences derived from the gene construct, for example nucleotide sequences derived from the left border sequence, right border sequence, origin of replication, recombinase-encoding or transposase-encoding sequence if used, or other vector-derived nucleotide sequences. Accordingly, a footprint is identifiable according to the nucleotide sequence of the recombination locus or transposon of the gene construct used, such as, for example, a sequence of nucleotides corresponding or complementary to a *lox* site or *frt* site.

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The term "cell cycle" means the cyclic biochemical and structural events associated with growth and with division of cells, and in particular with the regulation of the replication of DNA and mitosis. Cell cycle includes phases called: G0, Gap1 (G1), DNA synthesis (S), Gap2 (G2), and mitosis (M). Normally these four phases occur sequentially, however, the cell cycle also includes modified cycles wherein one or more phases are absent resulting in modified cell cycle such as endomitosis, acytokinesis, polyploidy, polyteny, and endoreduplication.

The term "cell cycle progression" refers to the process of passing through the different cell cycle phases. The term "cell cycle progression rate" accordingly refers to the speed at which said cell cycle phases are run through or the time spans required to complete said cell cycle phases.

With "two-hybrid assay" is meant an assay that is based on the observation that many eukaryotic transcription factors comprise two domains, a DNA-binding domain (DB) and an activation domain (AD) which, when physically separated (i.e. disruption of the covalent linkage) do not effectuate target gene expression. Two proteins able to interact physically with one of said proteins fused to DB and the other of said proteins fused to AD will re-unite the DB and AD domains of the transcription factor resulting in target gene expression. The target gene in the yeast two-hybrid assay is usually a reporter gene such as the β-galactosidase gene. Interaction between protein partners in the

yeast two-hybrid assay can thus be quantified by measuring the activity of the reporter gene product (Bartel and Fields 1997). Alternatively, a mammalian two-hybrid system can be used which includes e.g. a chimeric green fluorescent protein encoding reporter gene (Shioda *et al.*, 2000).

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Furthermore, folding simulations and computer redesign of structural motifs of the protein of the invention can be performed using appropriate computer programs (Olszewski, Proteins 25 (1996), 286-299; Hoffman, Comput. Appl. Biosci. 1 (1995), 675-679). Computer modeling of protein folding can be used for the conformational and energetic analysis of detailed peptide and protein models (Monge, J. Mol. Biol. 247 (1995), 995-1012; Renouf, Adv. Exp. Med. Biol. 376 (1995), 37-45). In particular, the appropriate programs can be used for the identification of interactive sites of the cytokinin oxidases, its ligands or other interacting proteins by computer assistant searches for complementary peptide sequences (Fassina, Immunomethods 5 (1994), 114-120). Further appropriate computer systems for the design of protein and peptides are described in the prior art, for example in Berry, Biochem. Soc. Trans. 22 (1994). 1033-1036; Wodak, Ann, N. Y. Acac. Sci. 501 (1987), 1-13; Pabo, Biochemistry 25 (1986), 5987-5991. The results obtained form the above-described computer analysis can be used for, e.g. the preparation of peptidomimetics of the protein of the invention or fragments thereof. Such pseudopeptide analogues of the natural amino acid sequence of the protein may very efficiently mimic the parent protein (Benkirane, J. Biol. Chem. 271 (1996), 33218-33224). For example, incorporation of easily available achiral Ω amino acid residues into a protein of the invention or a fragment thereof results in the substitution of amino bonds by polymethylene units of an aliphatic chain, thereby providing a convenient strategy for constructing a peptidomimetic (Banerjee, Biopolymers 39 (1996), 769-777). Superactive peptidomimetic analogues of small peptide hormones in other systems are described in the prior art (Zhang, Biochem. Biophys. Res. Commun. 224 (1996), 327-331). Appropriate peptidomimetics of the protein of the present invention can also be identified by the synthesis of peptidomimetic combinatorial libraries through successive amine alkylation and testing the resulting compounds, e.g., for their binding, kinase inhibitory and/or immunlogical properties. Methods for the generation and use of peptidomimetic combinatorial libraries are described in the prior art, for example in Ostresh, Methods in Enzymology 267 (1996), 220-234 and Dorner, Bioorg. Med. Chem. 4 (1996), 709-715.

Furthermore, a three-dimensional and/or crystallographic structure of the protein of the invention can be used for the design of peptidomimetic inhibitors of the biological activity of the protein of the invention (Rose, Biochemistry 35 (1996), 12933-12944; Ruterber, Bioorg. Med. Chem. 4 (1996), 1545-1558).

The compounds to be obtained or identified in the methods of the invention can be compounds that are able to bind to any of the nucleic acids, peptides or proteins of the invention. Other interesting compounds to be identified are compounds that modulate the expression of the genes or the proteins of the invention in such a way that either the expression of said gene or protein is enhanced or decreased by the action of said compound. Alternatively the compound can exert his action by enhancing or decreasing the activity of any of the proteins of the invention. Herein, preferred proteins are novel cytokinin oxidases.

Said compound or plurality of compounds may be comprised in, for example, samples, e.g., cell extracts from, e.g., plants, animals or microorganisms. Furthermore, said compound(s) may be known in the art but hitherto not known to be capable of suppressing or activating cell cycle interacting proteins. The reaction mixture may be a cell free extract of may comprise a cell or tissue culture. Suitable set ups for the method of the invention are known to the person skilled in the art and are, for example, generally described in Alberts et al., Molecular Biology of the Cell, third edition (1994), in particular Chapter 17. The plurality of compounds may be, e.g., added to the reaction mixture, culture medium or injected into the cell.

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If a sample containing a compound or a plurality of compounds is identified in the method of the invention, then it is either possible to isolate the compound form the original sample identified as containing the compound capable of acting as an agonist, or one can further subdivide the original sample, for example, if it consists of a plurality of different compounds, so as to reduce the number of different substances per sample and repeat the method with the subdivisions of the original sample. Depending on the complexity of the samples, the steps described above can be performed several times, preferably until the sample identified according to the method of the invention only comprises a limited number of or only one substance(s). Preferably said sample comprises substances or similar chemical and/or physical properties, and most preferably said substances are identical. Preferably, the compound identified according to the above-described method or its derivative is further formulated in a form suitable for the application in plant breeding or plant cell and tissue culture.

The term "early vigor" refers to the ability of a plant to grow rapidly during early development, and relates to the successful establishment, after germination, of a well-developed root system and a well-developed photosynthetic apparatus.

The term "resistance to lodging" or "standability" refers to the ability of a plant to fix itself to the soil. For plants with an erect or semi-erect growth habit this term also refers to the ability to maintain an upright position under adverse (environmental) conditions. This trait relates to the size, depth and morphology of the root system.

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As used herein, tblastn refers to an alignment tool that is part of the BLAST (Basic Local Alignment Search Tool) family of programs (http://www.ncbi.nlm.nih.gov/BLAST/). BLAST aims to identify regions of optimal local alignment, i.e. the alignment of some portion of two nucleic acid or protein sequences, to detect relationships among sequences which share only isolated regions of similarity (Altschul et al., 1990). In the present invention, tblastn of the BLAST 2.0 suite of programs was used to compare the maize cytokinin oxidase protein sequence against a nucleotide sequence database dynamically translated in all reading frames (Altschul et al., Nucleic Acids Res. 25: 3389-3402 (1997)).

The following examples and figures are given by means of illustration of the present invention and are in no way limiting. The contents of all references included in this application are incorporated by reference.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1. Schematic representation of plant cytokinin oxidase genes.

Shown are the structures of different cytokinin oxidase genes isolated from maize (ZmCKX1, accession number AF044603, Biochem. Biophys. Res. Com. 255:328-333, 1999) and Arabidopsis (AtCKX1 to AtCKX4). Exons are denominated with 'E' and represented by shaded boxes. Introns are represented by white boxes. Further indicated are the gene sizes (in kb, on top of each structure), the gene accession numbers (under the names) and a size bar representing 0.5 kb.

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Figure 2. Alignment of plant cytokinin oxidase amino acid sequences.

The amino acid sequences from cytokinin oxidases from maize (ZmCKX1) and Arabidopsis (AtCKX1 to AtCKX4) are aligned. Identical amino acid residues are marked by a black box, similar amino acid residues are in a grey box. Amino acid similarity groups: (M,I,L,V), (F,W,Y), (G,A), (S,T), (R,K,H), (E,D), (N,Q),

Figure 3. Northern blot analysis of *AtCKX1*-expressing tobacco and *Arabidopsis* plants.

- (A) Northern blot analysis of constitutively expressing tobacco plants (lanes 1-8) compared to wild type SNN tobacco (lane 9)
- (B) Comparison of tetracycline-induced gene expression in leaves after 12h of induction with a constitutively expressing clone. Lanes 2-9, leaves of four different *CKX1*-W38TetR clones (+,-, with or without tetracycline treatment), lane 1, constitutively expressing 35S::CKX1 clone.
- (C) Northern blot analysis of Arabidopsis plants constitutively expressing CKX1 gene. Lanes 2-4, three different constitutively expressing 35S::CKX1 clones compared to wild type Arabidopsis plant (lane 1).

Figure 4: Growth characteristics of 35S::CKX1 transgenic Arabidopsis plants.

- (A) Two wild type seedlings (left) compared to two 35S::CKX1 expressing seedlings (right). Note the increased formation of adventitious roots and increased root branching in the trangenic seedlings. Pictures were taken 14 days after germination. Plants were grown in vitro on MS medium in petri dishes in a vertical position.
 - (B) Like A, but roots stained with toluidine blue.

- (C) Top view of a petri dish with 35S::CKX1 transgenic seedlings three weeks after germination.
- (D) A 35S::CKX1 transgenic plants grown in liquid culture. Roots of wild type seedlings grow poorly under these conditions (not shown).
- (E) Transformants (T0) that express the 35S::CKX1 gene (three plants on the right), a wild type plant is shown on the left.
 - (F) Phenotype of T1 plants grown in soil. Wild type plant (left) compared to two 35S::CKX1 trangenic plants.

Figure 5: Phenotype of CKX2 overexpressing Arabidopsis plants.

T1 generation of 35S::CKX2 expressing Arabidopsis plants (two plants on the right) compared to wild type (plant on the left).

Figure 6. Northern blot analysis of AtCKX2-expressing tobacco and Arabidopsis plants.

- (A) Northern blot analysis of constitutively expressing tobacco plants (lanes 1-7) compared to wild type SNN tobacco (lane 8)
- (B) Northern blot analysis of Arabidopsis plants constitutively expressing *CKX2* gene. Lanes 2-8, seven different consitutively expressing *35S::CKX2* clones compared to wild type Arabidopsis plant (lane 1).

Figure 7. Shoot phenotype of AtCKX1 and AtCKX2 expressing tobacco plants.

(A) Top view of six week old plants.

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- (B) Tobacco plants at the flowering stage.
- (C) Kinetics of stem elongation. Arrows mark the onset of flowering. Age of plants (days after germination) and leaf number at that stage are indicated above the arrows. Bars indicate SD; n = 12.
 - (D) Number of leaves (n = 12) formed between day 68 and day 100 after germination and final surface area of these leaves (100% of wild type is 3646 ±144 cm²; n = 3).
- (E) Comparison of leaf size and senescence. Leaves were from nodes number 4, 9, 12, 16 and 20 from the top (from left to right).

Figure 8. Root phenotype of AtCKX expressing transgenic tobacco plants.

(A) Seedlings 17 days after germination.

- (B) Root system of soil grown plants at the flowering stage.
- (C) Root length, number of lateral roots (LR) and adventitious roots (AR) on day 10 after germination.
- (D) Dose-response curve of root growth inhibition by exogenous cytokinin. Bars indicate \pm SD; n = 30.

Figure 9: Growth of axillary shoot meristems in 35S::CKX1 expressing tobacco plants.

- Figure 10: Histology of shoot meristems, leaves and root meristems of *AtCKX1* overexpressing tobacco plants versus wild type (WT) tobacco.
 - (A) Longitudinal median section through the vegetative shoot apical meristem. P, leaf primordia.
 - (B) Vascular tissue in second order veins of leaves. X, xylem, PH, a phloem bundle.
- 15 (C) Cross sections of fully developed leaves.

- (D) Scanning electron microscopy of the upper leaf epidermis.
- (E) Root apices stained with DAPI. RM, root meristem.
- (F) Longitudinal median sections of root meristems ten days after germination. RC, root cap; PM, promeristem.
- (G) Transverse root sections 10 mm from the apex. E, epidermis, C1-C4, cortical cell layer, X, xylem, PH, phloem. Bars are 100 μm.

EXAMPLES

Example 1. Identification of candidate cytokinin oxidase encoding genes from Arabidopsis thaliana

Six different genes are identified from Arabidopsis thaliana that bear sequence similarity 5 to a cytokinin oxidase gene from maize (Morris et al., Biochem Biophys Res Comm 255:328-333, 1999; Houda-Herin et al. Plant J 17:615-626; WO 99/06571). These genes are found by screening 6-frame translations of nucleotide sequences from public genomic databases with the maize protein sequence, employing tblastn program. These sequences are designated as Arabidopsis thaliana cytokinin oxidase-like genes or 10 AtCKX. They are arbitrarily numbered as AtCKX1 to AtCKX6. The below list summarizes the information on these genes. The predicted ORF borders and protein sequences are indicative, in order to illustrate by approximation the protein sequence divergence between the Arabidopsis and maize cytokinin oxidases, as well as amongst the different Arabidopsis cytokinin oxidases. The ORF borders and protein sequences shown should not be taken as conclusive evidence for the mode of action of these AtCKX genes. The % identity/similarity values that are indicated amongst the Arabidopsis genes represent the lowest and highest values found with all combinations.

A. Gene name: AtCKX1 (Arabidopsis thaliana cytokinin oxidase-like protein 1, SEQ ID 20 NO1)

Location in database (accession number, location on bac): AC002510, Arabidopsis thaliana chromosome II section 225 of 255 of the complete sequence. Sequence from clones T32G6.

ORF predicted in the database:

15517..16183, 16415..16542, 16631..16891, 16995..17257, 17344..17752

Predicted protein sequence: SEQ ID NO 2 30

Homologies

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% identity with Z. mays gene:

31,5% (Dnastar/MegAlign - Clustal method)

% similarity with Z. mays protein: 35

32,2% (Dnastar/MegAlign - Clustal method)

% identity with other Arabidopsis genes (range):

38,2% (AtCKX2) - 54,1% (AtCKX6) (Dnastar/MegAlign - Clustal method)

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% similarity with other Arabidopsis proteins (range):

37,1% (AtCKX2) - 58,1% (AtCKX6) (Dnastar/MegAlign - Clustal method)

B. Gene name: AtCKX2 (Arabidopsis thaliana cytokinin oxidase-like protein 2, SEQ ID NO3)

Location in database (accession number, location on bac): AC005917, Arabidopsis thaliana chromosome II section 113 of 255 of the complete sequence. Sequence from clones F27F23, F3P11.

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ORF predicted in the database:

complement, 40721..41012, 41054..41364, 41513..41770, 42535..42662, 43153..43711

Please note: The cDNA sequence is different than annotated in the database. Based on the cDNA sequence and using the gene prediction program NetPlantGene (http://www.cbs.dtu.dk/services/NetGene2/) the ORF predicted in the database is revised:

complement, 40721..41012, 41095..41364, 41513..41770, 42535..42662, 43153..43711

25 Predicted protein sequence, based on corrected ORF: SEQ ID NO 4

Homologies

% identity with Z. mays gene:

38,4% (Dnastar/MegAlign - Clustal method)

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% similarity with Z. mays protein:

37,5% (Dnastar/MegAlign - Clustal method)

% identity with other Arabidopsis genes (range):

34,9% (AtCKX6) - 64,5% (AtCKX4) (Dnastar/MegAlign - Clustal method)

% similarity with other Arabidopsis proteins (range):

36,5% (AtCKX6) - 66,1% (AtCKX4) (Dnastar/MegAlign - Clustal method)

C. Gene name: AtCKX3 (Arabidopsis thaliana cytokinin oxidase-like protein 3, SEQ ID NO 5)

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Location in database (accession number, location on bac): AB024035, Arabidopsis thaliana genomic DNA, chromosome 5, P1 clone: MHM17, complete sequence.

No prediction of the ORF in the database.

The gene is identified by several gene prediction programs including GRAIL (ftp: 10 //arthur.epm.ornl.gov/pub/xgrail), Genscan (http://CCR-081.mit.edu/GENSCAN .html) and NetPlantGene (http://www.cbs.dtu.dk/services/NetGene2/):

complement, 29415..29718, 29813..30081, 30183..30443, 30529..30656, 32107..32716

Predicted protein sequence, based on own ORF prediction: SEQ ID NO 6

Homologies

% identity with Z. mays gene:

38,7% (Dnastar/MegAlign - Clustal method)

% similarity with Z. mays protein:

39,2% (Dnastar/MegAlign - Clustal method)

% identity with other Arabidopsis genes (range): 25

38,8% (AtCKX6) - 51,0% (AtCKX2) (Dnastar/MegAlign - Clustal method)

% similarity with other Arabidopsis proteins (range):

39,9% (AtCKX6) - 46,7% (AtCKX2) (Dnastar/MegAlign - Clustal method)

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D. Gene name: AtCKX4 (Arabidopsis thaliana cytokinin oxidase-like protein 4, SEQ ID NO 7)

Location in database (accession number, location on bac):

- 1) AL079344, Arabidopsis thaliana DNA chromosome 4, BAC clone T16L4 (ESSA 35 project)
 - 2) AL161575, Arabidopsis thaliana DNA chromosome 4, contig fragment No. 71.

ORF predicted in the database:

- 1) 76187..76814, 77189..77316, 77823..78080, 78318..78586, 78677..78968
- 2) 101002..101629, 102004..102131, 102638..102895, 103133..103401,
- 5 103492..103783

Predicted protein sequence: SEQ ID NO 8

Homologies

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10 % identity with Z. mays gene:

41,0% (Dnastar/MegAlign - Clustal method)

% similarity with Z. mays protein:

41,0% (Dnastar/MegAlign - Clustal method)

% identity with other *Arabidopsis* genes (range):

35,2% (AtCKX6) - 64,5% (AtCKX2) (Dnastar/MegAlign - Clustal method)

% similarity with other Arabidopsis proteins (range):

35,1% (AtCKX6) - 66,1% (AtCKX2) (Dnastar/MegAlign - Clustal method)

E. Gene name: AtCKX5 (Arabidopsis thaliana cytokinin oxidase-like protein 5, SEQ ID NO 9)

Location in database (accession number, location on bac): AC023754, F1B16, complete sequence, chromosome 1

No prediction of the ORF in the database.

The gene can be identified by several gene prediction programs including GRAIL (ftp://arthur.epm.ornl.gov/pub/xgrail), Genscan (http://cch.ornl.gov/pub/xgrail), Genscan (http://cch.ntm)

43756..44347, 44435..44562, 44700..44966, 45493..45755, 46200..46560

35 Predicted protein sequence: SEQ ID NO 10

Homologies

% identity with Z. mays gene:

39,1% (Dnastar/MegAlign - Clustal method)

% similarity with Z. mays protein:

36,6% (Dnastar/MegAlign - Clustal method)

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% identity with other Arabidopsis genes (range):

40,1% (AtCKX2) - 44,0% (AtCKX3) (Dnastar/MegAlign - Clustal method)

% similarity with other Arabidopsis proteins (range):

41,6% (AtCKX4) - 46,4% (AtCKX6) (Dnastar/MegAlign - Clustal method)

F. Gene name: AtCKX6 (Arabidopsis thaliana cytokinin oxidase-like protein 6, SEQ ID NO 11)

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Location in database (accession number, location on bac): AL163818, *Arabidopsis thaliana* DNA chromosome 3, P1 clone MAA21 (ESSA project).

ORF predicted in the database:

20 46630..47215, 47343..47470, 47591..47806, 47899..48161, 48244..48565

Predicted protein sequence: SEQ ID NO 12

Homologies

% identity with Z. mays gene:

37,3% (Dnastar/MegAlign - Clustal method)

% similarity with Z. mays protein:

36,1% (Dnastar/MegAlign - Clustal method)

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% identity with other Arabidopsis genes (range):

34,9% (AtCKX2) - 54,1% (AtCKX1) (Dnastar/MegAlign - Clustal method)

% similarity with other Arabidopsis proteins (range):

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35,1% (AtCKX4) – 58,1% (AtCKX1) (Dnastar/MegAlign - Clustal method) Genes AtCKX3 and AtCKX5 are not annotated as putative cytokinin oxidases in the database and ORFs for these genes are not given. Furthermore, the ORF (and

consequently the protein structures) predicted for AtCKX2 is different from our own prediction.

A comparison of the gene structure of the *Arabidopsis AtCKX* genes 1 to 4 and the maize *CKX* gene is shown in Fig 1.

The predicted proteins encoded by the *Arabidopsis AtCKX* genes show between 32% and 41% sequence similarity with the maize protein, while they show between 35% and 66% sequence similarity to each other. Because of this reduced sequence conservation, it is not clear *a priori* whether the *Arabidopsis AtCKX* genes encode proteins with cytokinin oxidase activity. An alignment of the *Arabidopsis AtCKX* predicted proteins 1 to 4 and the maize *CKX* gene is shown in Fig 2.

Example 2. Transgenic plants overexpressing AtCKX1 show increased cytokinin oxidase activity and altered plant morphology

15 1. Description of the cloning process

The following primers are used to PCR amplify the AtCKX1 gene from *Arabidopsis* thaliana, accession Columbia:

Sequence of 5' primer: cggtcgacATGGGATTGACCTCATCCTTACG (SEQ ID NO:13)
Sequence of 3' primer: gcgtcgacTTATACAGTTCTAGGTTTCGGCAGTAT (SEQ ID NO:

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A 2235-bp PCR fragment, amplified by these primers, is inserted in the Sal I site of pUC19. The insert is sequenced to confirm that the PCR amplification has not introduced mutations. The Sall/Sall fragment of this vector is subcloned in the Sall site downstream of a modified CaMV 35S promoter (carrying three tetracycline operator sequences) in the binary vector pBinHyg-Tx (Gatz et al., 1992). The resulting construct is introduced into tobacco and *Arabidopsis thaliana* through *Agrobacterium*-mediated transformation, using standard transformation protocols.

2. Molecular analysis of the transgenic lines

Several transgenic lines are identified that synthesize the *AtCKX1* transcript at high levels (Fig 3). Transgenic lines expressing *AtCKX1* transcript also show increased cytokinin oxidase activity. This is exemplified for 2 tobacco and 2 Arabidopsis lines in

Table 6. This result proves that the AtCKX1 gene encodes a protein with cytokinin oxidase activity.

Table 6. Cytokinin oxidase activity in AtCKX1 transgenic plant tissues

Leaf sample		
Plant species	Plant line	Cytokinin oxidase activity (nmol Ade/mg protein.h)
Arabidopsis	Col-0 wild-type	0.009
	CKX1-11	0.024
	CKX1-22	0.026
	CKX1-22	0.027
Tobacco	SNN wild-type	0.004
	CKX1-SNN-8	0.016
	CKX1-SNN-28	0.021

3. Phenotypic description of the transgenic lines

In Arabidopsis (see also Fig 4):

onset of germination same as for wt

- the number of adventitious roots early after germination is enhanced; the growth of aerial organs (hypocotyl, cotyledons and leaves) is delayed and reduced resulting in a dwarfed phenotype
- the number of leaves in rosette is increased

- the life cycle is longer compared to wt

In tobacco (see also Fig 7 to 10):

The plants have a dwarfed phenotype, with reduced apical dominance and increased root production.

Five categories of phenotype:

- 1) strong 2 clones
- 2) intermediate 3 clones
- 3) weak 4 clones

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- 4) tall plants (as WT) with large inflorescence 5 clones
- 5) similar to wt, 9 clones

Height (see Fig. 7)

- 5
- wt: between 100-150 cm
- weak: approximately 75 cm
- intermediate: appr. 40-45 cm (main stem app. 25 cm but overgrown by side branches.
- strong: appr. 10 cm

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Roots (see Fig. 8)

In vitro grown plants highly expressing the gene are easily distinguishable from the WT by their ability to form more roots which are thicker (stronger) as well as by forming aerial roots along the stem.

- 15 Roots in soil (4-4,5 month after transfer to the green house):
 - strong: good root system (as WT) despite the strong reduction of aerial parts
 - intermediate: large increase in root system (biomass)
 - weak: also increase in root biomass compared to WT

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Internodes distance

- wt: 5th internode below inflorescence app. 5 cm, 9th internode about 2 cm long
- intermediate class: 5th internode below inflorescence about 2.5 cm, 9th internode app. 0,5 cm long

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Seed capsules

Expressing plants have in general less capsules, which are of similar size as wt (even in strong expressors)

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- strong: 2-4 capsules
- intermediate: about 25
- wt: more than 60

Flower

Size is not reduced. The first flowers are aborted in the transgenic expressors.

Leaves (see Figure 7)

The shape of leaves of *AtCKX1* transgenic expressors is lanceolate (longer and narrow). The width-to-length ratio of mature leaves was reduced from 1:2 in wild type plants to 1:3 in *AtCKX1* transgenics. A prominent difference was also noted for progression of leaf senescence. In wt tobacco, leaf senescence starts in the most basal leaves and leads to a uniform reduction of leaf pigment. By contrast, ageing leaves of strongly expressing *AtCKX1* plants stay green along the leaf veins and turn yellow in the intercostal regions, indicating altered leaf senescence. The texture of older leaves is more rigid.

10 Stem

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More side branches are formed indicating reduced apical dominance compared to WT plants (see Figure 9). The side branches overgrow the main stem, reaching a height of 40-45 cm for intermediate *AtCKX1* expressors. Even secondary branches appear.

Example 3. Transgenic plants overexpressing AtCKX2 show increased cytokinin oxidase activity and altered plant morphology

1. Description of the cloning process

The following primers are used to PCR amplify the AtCKX2 gene from *Arabidopsis* thaliana, accession Columbia:

Sequence of 5' primer: gcggtaccAGAGAGAGAGAAACATAAACAAATGGC (SEQ ID NO:15)

Sequence of 3' primer: gcggtaccCAATTTTACTTCCACCAAAATGC (SEQ ID NO:16)

A 3104-bp PCR fragment, amplified by these primers, is inserted in the KpnI site of pUC19. The insert is sequenced to check that no differences to the published sequence are introduced by the PCR procedure. The KpnI/KpnI fragment of this vector is subcloned in the KpnI site downstream of a modified CaMV 35S promoter (carrying three tetracycline operator sequences) in the binary vector pBinHyg-Tx (Gatz et al., 1992). The resulting construct is introduced into tobacco and Arabidopsis thaliana through Agrobacterium-mediated transformation, using standard transformation protocols.

2. Molecular analysis of the transgenic lines

Several transgenic lines are identified that synthesize the *AtCKX2* transcript at high levels (Fig 6). Transgenic lines expressing *AtCKX2* transcript also show increased cytokinin oxidase activity. This is exemplified for 2 tobacco and 3 Arabidopsis lines in Table 7. This result proves that the *AtCKX2* gene encodes a protein with cytokinin oxidase activity.

Table 7. Cytokinin oxidase activity in AtCKX2 transgenic plant tissues

San	nple	7			
Plant species and tissue	Plant line	Cytokinin oxidase activity (nmol Ade/mg protein.h)			
Arabidopsis callus	Col-0 wild-type	0.037			
	CKX2-15	0.351			
	CKX2-17	0.380			
	CKX2-55	0.265			
Tobacco leaves	SNN wild-type	0.009			
	CKX2-SNN-18	0.091			
	CKX2-SNN-19	0.091			

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3. Phenotypic description of the transgenic lines

In Arabidopsis (see Fig 5):

Basically similar to CKX1 (no quantitative comparisons with AtCKX1 available)

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In tobacco (see Fig 7 to 10):

Three categories of phenotype:

- 1) strong 15 clones (similar to intermediate phenotype of AtCKX1)
- 2) weak 6 clones
- 3) others similar to wt plants, 7 clones

Roots

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In vitro grown plants highly expressing the gene are easily distinguishable from wt plants by their ability to form more roots which are thicker (stronger) as well as by forming aerial roots along the stem.

- 5 Roots in soil (4-4,5 month after transfer to the green house):
 - strong: very significant increase in root biomass for strongest expressing clones
 - weak: also increase in root biomass compared to wt
- The observations concerning plant height, internode distance, branching, seed set, leaf form and yellowing are similar as for *AtCKX1* transgenics with some generally minor quantitative differences.

Interestingly, the increase in cytokinin oxidase activity and in root production is more pronounced in *AtCKX2* than in *AtCKX1* expressing tobacco lines. This is also illustrated by the lower cytokinin content of *AtCKX2* plants versus *AtCKX1* plants (see Table 9 below). These results indicate that *AtCKX2* may be a preferential tool for the embodiments of this patent application as compared to *AtCKX1*.

Example 4. Transgenic plants overexpressing AtCKX3 show altered plant morphology

1. Description of the cloning process

The following primers are used to PCR amplify the *AtCKX3* gene from *Arabidopsis* thaliana, accession Columbia:

Sequence of 5´ primer: gcggtaccTTCATTGATAAGAATCAAGCTATTCA (SEQ ID NO:17)

Sequence of 3' primer: gcggtaccCAAAGTGGTGAGAACGACTAACA (SEQ ID NO:18)

A 3397-bp PCR fragment, produced by this PCR amplification, is inserted in the Kpnl site of pBluescript. The insert is sequenced to confirm that the PCR product has no sequence changes as compared to the gene. The Kpnl/Kpnl fragment of this vector is subcloned in the Kpnl site downstream of a modified CaMV 35S promoter (carrying three tetracycline operator sequences) in the binary vector pBinHyg-Tx (Gatz et al., 1992).

2. Plant phenotypic analysis

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The phenotypes generated by overexpression of the *AtCKX3* gene in tobacco and *Arabidopsis* are basically similar as those of *AtCKX1* and *AtCKX2* expressing plants, i.e. enhanced rooting and dwarfing.

Example 5. Transgenic plants overexpressing AtCKX4 show altered plant morphology

1. Description of the cloning process

The following primers are used to PCR amplify the AtCKX4 gene from *Arabidopsis* thaliana, accession Columbia:

Sequence of 5' primer: gcggtaccCCCATTAACCTACCCGTTTG (SEQ ID NO:19)
Sequence of 3' primer: gcggtaccAGACGATGAACGTACTTGTCTGTA (SEQ ID NO:20)

A 2890-bp PCR fragment, produced by this PCR amplification, is inserted in the Kpnl site of pBluescript. The insert is sequenced to confirm that the PCR product has no sequence changes as compared to the gene. The Kpnl/Kpnl fragment of this vector is subcloned in the Kpnl site downstream of a modified CaMV 35S promoter (carrying three tetracycline operator sequences) in the binary vector pBinHyg-Tx (Gatz et al., 1992).

2. Molecular analysis of the transgenic lines

Several transgenic lines are identified that show increased cytokinin oxidase activity. This is exemplified for 3 Arabidopsis lines in Table 8. This result proves that the AtCKX4 gene encodes a protein with cytokinin oxidase activity.

Table 8. Cytokinin oxidase activity in AtCKX4 transgenic plant tissues

San	nple			
Plant species and tissue	Plant line	Cytokinin oxidase activity (nmol Ade/mg protein.h)		
Arabidopsis callus	Col-0 wild-type	0.037		
	CKX4-37	0.244		
· · · · · · · · · · · · · · · · · · ·	CKX4-40	0.258		
	CKX4-41	0.320		

3. Plant phenotypic analysis

The phenotypes generated by overexpression of the *AtCKX4* gene in tobacco and *Arabidopsis* are basically similar as those of *AtCKX1* and *AtCKX2* expressing plants, i.e. enhanced rooting, reduced apical dominance, dwarfing and yellowing of intercostal regions in older leaves of tobacco. An additional phenotype in tobacco are lanceolate leaves (altered length-to-wide ratio).

Example 6. Cloning of the AtCKX5 gene

The following primers are used to PCR amplify the AtCKX5 gene from *Arabidopsis* thaliana, accession Columbia:

Sequence of 5' primer: ggggtaccGTGAAATGACGTCAAGCTTTCTT (SEQ ID NO:21)
Sequence of 3' primer: ggggtaccCTTTCCTCTTGGTTTTGTCCTGT (SEQ ID NO:22)

A 2833-bp PCR fragment, produced by this PCR amplification, is inserted as a blunt-end product in pCR-Blunt II-TOPO cloning vector (Invitrogen).

Example 7. Cloning of the AtCKX6 gene

The following primers are used to PCR amplify the AtCKX6 gene from *Arabidopsis* thaliana, accession Columbia:

Sequence of 5' primer: gctctagaTCAGGAAAAGAACCATGCTTATAG (SEQ ID NO:23)
Sequence of 3' primer: gctctagaTCATGAGTATGAGACTGCCTTTTG (SEQ ID NO:24)

A 1949-bp PCR fragment, produced by this PCR amplification, is inserted as a blunt-end product in pCR-Blunt II-TOPO cloning vector (Invitrogen).

Example 8. Expression of the AtCKX2 gene under a root-specific promoter in transgenic tobacco leads to increased root production

The AtCKX2 gene (see example 3) is cloned under control of the WRKY6 promoter of Arabidopsis, which is a promoter that drives root-specific expression.

Transgenic plants expressing the *AtCKX2* gene specifically in the roots show increased root production without negatively affecting growth and development of the aerial parts of the plant.

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Example 9. Tobacco seedling growth test demonstrate early vigor of AtCKX transgenics

Seeds of *AtCKX1*-, *AtCKX2*-overexpressing and wt tobacco were sown in vitro, brought to culture room after 4 days (after cold treatment), and germinated after 6 days.

Observation on seedling growth were made 13 days after sowing:

Wild type:

just two cotyledons, tiny leaves visible;

root length 9-11 mm, no side root, no adventitious root

AtCKX1:

clone 50 (strongest phenotype in green house): root length 18-21 mm, 1-2 side branches (sb), 1-2 adventitious roots (ar)

15 clone 8: root length 13-16 mm, 1 ar, 1 sb

clone 2: root length 17-20 mm, 1-2 ar, 1-2 sb

AtCKX2:

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clone 2: root length 14-18 mm

clone 38: root length 16-19 mm

clone 40: root length 15-20 mm

clone 2 (very low expresser): root length 10 mm

formation of more adventitious roots also evident for AtCKX2 transgenics

25 AtCKX1 and AtCKX2 plants, general observations:

Longer primary root (up to 100%!) which has already branched (1-2 side branches), more adventitious roots (1-2). Primary true leaves are advanced, cotyledons are somewhat more expanded. These results show that overexpression of cytokinin oxidase enhances the growth and development of both the main root and the adventitious roots, resulting in early vigor.

Example 10. Histological analysis of altered plant morphology in *AtCKX1* overexpressing tobacco plants

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Microscopic analysis of different tissues revealed that the morphological changes in AtCKX transgenics are reflected by distinct changes in cell number and rate of cell formation (see Figure 10). The shoot apical meristem (SAM) of AtCKX1 transgenics is smaller than in wild type and fewer cells occupy the space between the central zone and the peripheral zone of lateral organ fromation, but the cells are of the same size (Figure 10 A). The reduced cell number and size of the SAM as a consequence of a reduced cytokinin content indicates that cytokinins have a role in the control of SAM proliferation. No obvious changes in the differentiation pattern occurred, suggesting that the spatial organization of the differentiation zones in the SAM is largely independent from cellnumber and from the local cytokinin concentration. The overall tissue pattern of leaves in cytokinin oxidase overexpressers is unchanged. However, the size of the phloem and xylem is significantly reduced (Figure 10 B). By contrast, the average cell size of leaf parenchyma and epidermal cells is increased four- to fivefold (Figure 10 C, D). New cells of AtCKX1 transgenics are formed at 3-4% of the rate of wild type leaves and final leaf cell number is estimated to be in the range of 5-6% of wild type. This indicates an absolute requirement for cytokinins in leaves to maintain the cell division cycle. Neither cell size nor cell form of floral organs was altered and seed yield per capsule was similar in wild type and AtCKX transgenic plants. The cell population of root meristems of AtCKX1 transgenic plants was enlarged approximately 4-fold and the cell numbers in both the central and lateral columnella were enhanced (Figure 10 E, F). The final root diameter was increased by 60% due to an increased diameter of all types of root cells. The radial root patterns was identical in wild type and transgenics, with the exception that frequently a fourth layer of cortex cells was noted in transgenic roots (Figure 10 G). The increased cell number and the slightly reduced cell length indicates that the enhanced root growth is due to an increased number of cycling cells rather than increased cell growth. In the presence of lowered cytokinin content, root meristem cells must undergo additional rounds of mitosis before they leave the meristem and start to elongate. We conclude that the exit from the meristem is regulated by a mechanism that is sensitive to cytokinins. Apparently, cytokinins have a negative regulatory role in the root meristem and wild type cytokinin concentrations are inhibitory to the development of a maximal root system. Therefore, reducing the level of active cytokinins by overexpressing cytokinin oxidases stimulates root development which results in an increase in the size of the root with more lateral and adventitious roots as compared to wt plants.

Example 11. The endogenous concentration of different cytokinin metabolites is significantly reduced in *AtCKX* expressing transgenic tobacco seedlings.

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Among the 16 different cytokinin metabolites that were measured, the greatest change occurred in the iP-type cytokinins in *AtCKX2* overexpressers (Table 9). Smaller alterations were noted for Z-type cytokinins, which could be due to a different accessibility of the substrate or a lower substrate specificity of the protein. The total content of iP and Z metabolites in individual transgenic clones was between 31% and 63% of wild type. The cytokinin reserve pool of *O*-glucosides was also lowered in the transgenics (Table 9). The concentration of *N*-glucosides and DHZ-type cytokinins was very low and was not or only marginally, altered in transgenic seedlings (data not shown).

Table 9. Cytokinin content of *AtCKX* transgenic plants. Cytokinin extraction, immunopurification, HPLC separation and quantification by ELISA methods was carried out as described by Faiss et al., 1997. Three independently pooled samples of approximately 100 two week old seedlings (2.5 g per sample) were analyzed for each clone. Concentrations are in pmol x g fresh weight⁻¹. Abbreviations: iP, N⁶-(Δ²isopentenyl)adenine; iPR, N⁶-(Δ²isopentenyl)adenine riboside; iPRP, N⁶-(Δ²isopentenyl)adenine riboside 5΄-monophosphate; Z, *trans*-zeatin; ZR, zeatin riboside; ZRP, zeatin riboside 5΄-monophosphate; ZOG, zeatin *O*-glucoside; ZROG, zeatin riboside *O*-glucoside.

Line Cytokinin	Wild-type	AtCKX1-2		AtCKX1-28		AtCKX2-38		AtCKX2-40	
	Concen-	Concen-	% of						
metabolite	tration	tration	WT	tration	WT	tration	WT	tration	WT
iP	5,90 ± 1,80	4,76 ± 0,82	81	4,94 ± 2,62	84	1,82 ± 0,44	31	$2,85 \pm 0,62$	48
iPR	$2,36 \pm 0,74$	1,53 ± 0,14	65	$0,75 \pm 0,27$	32	$0,55 \pm 0,39$	23	$0,89 \pm 0,07$	38
iPRP	$3,32 \pm 0,73$	0,87 ± 0,26	26	$1,12 \pm 0,13$	34	$0,80 \pm 0,48$	24	1,68 ± 0,45	51
Z	$0,24 \pm 0,06$	$0,17 \pm 0,02$	71	$0,22 \pm 0,03$	92	0,21 ± 0,06	88	0,22 ± 0,02	92
ZR	$0,60 \pm 0,13$	$0,32 \pm 0,12$	53	0,34 ± 0,03	57	0,34 ± 0,15	57	0,32 ± 0,05	53
ZRP	$0,39 \pm 0,17$	$0,42 \pm 0,11$	107	$0,28 \pm 0,15$	72	0,06 ± 0,01	15	$0,17 \pm 0,06$	44
zog	$0,46 \pm 0,20$	$0,32 \pm 0,09$	70	0,26 ± 0,13	57	0,20 ± 0,07	43	0,12 ± 0,02	26
ZROG	$0,48 \pm 0,17$	$0,30 \pm 0,06$	63	$0,47 \pm 0,02$	98	0,23 ± 0,05	48	$0,30 \pm 0,13$	63
Total	13,75	8,69	63	8,38	61	4,21	31	6,55	48

Example 12. Suppression of a AtCKX gene under a senescence-induced promoter in transgenic plants leads to delayed leaf senescence and enhanced seed yield.

A chimeric gene construct derived from a *AtCKX* gene (see example 3) and designed to suppress expression of endogenous cytokinin oxidase gene(s) is cloned under control of a senescence-induced promoter. For example, promoters derived from senescence-associated genes (SAG) such as the SAG12 promoter can be used (Quirino et al., 2000). Transgenic plants suppressing endogenous cytokinin oxidase gene(s) specifically in senescing leaves show delayed leaf senescence and higher seed yield without negatively affecting the morphology and growth and development of the plant.

Example 13. Overexpression of a AtCKX gene in the female reproductive organs leads to parthenocaropic fruit development

The open reading frame of an *AtCKX* gene is cloned under control of a promoter that confers overexpression in the female reproductive organs such as for example the DefH9 promoter from *Antirrhinum majus* or one of its homologues which has a high expression specificity in the placenta and ovules. Transgenic plants with enhanced cytokinin oxidase activity in these tissues show parthenocarpic fruit development.

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CLAIMS

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- A method for stimulating root growth and/or enhancing the formation of lateral and adventitious roots and/or altering root geotropism comprising expression of a plant cytokinin oxidase or comprising expression of another protein that reduces the level of active cytokinins in plants or plant parts.
- 2. An isolated nucleic acid encoding a novel plant protein having cytokinin oxidase activity or encoding an immunologically active and/or functional fragment of such a protein selected from the group consisting of:
- (a) nucleic acid sequences comprising a DNA sequence as given in any of SEQ ID NOs 1, 3, 5, 7, 9 or 11, or the complement thereof,
 - (b) nucleic acid sequences comprising the RNA sequences corresponding to any of SEQ ID NOs 1, 3, 5, 7, 9 or 11, or the complement thereof,
 - (c) nucleic acid sequences hybridizing to the nucleotide sequence as defined in (a) or (b),
 - (d) nucleic acid sequences encoding a protein with an amino acid sequence which is at least 35% similar to the amino acid sequence as given in SEQ ID NO 2,
 - (e) nucleic acid sequences encoding a protein with an amino acid sequence which is at least 40% similar to the amino acid sequence as given in SEQ ID NO 4.
 - (f) nucleic acid sequences encoding a protein with an amino acid sequence which is at least 42% similar to the amino acid sequence as given in SEQ ID NO 6,
 - (g) nucleic acid sequences encoding a protein with an amino acid sequence which is at least 44% similar to the amino acid sequence as given in SEQ ID NO 8,
 - (h) nucleic acid sequences encoding a protein with an amino acid sequence which is at least 40% similar to the amino acid sequence as given in SEQ ID NO 10,
 - (i) nucleic acid sequences encoding a protein with an amino acid sequence which is at least 40% similar to the amino acid sequence as given in SEQ ID NO 12,
 - (j) nucleic acid sequences encoding a protein comprising the amino acid sequence as given in any of SEQ ID NOs 2, 4, 6, 8, 10 or 12,
 - (k) nucleic acid sequences which are degenerated as a result of the genetic code to a nucleotide sequence of a nucleic acid as given in any of SEQ ID NOs 1, 3, 5, 7, 9 or 11, or as defined in (a) to (i),

- (I) nucleic acid sequences which are diverging due to the differences in codon usage between the organisms to a nucleotide sequence encoding a protein as given in any of SEQ ID NOs 2, 4, 6, 8, 10, 12 or as defined in (a) to (j),
- (m) nucleic acid sequences which are diverging due to the differences between alleles encoding a protein as given in SEQ ID NOs 2, 4, 6, 8, 10 or 12, or as defined in (a) to (j),
- (n) nucleic acid sequences encoding an immunologically active and/or functional fragment of a protein encoded by a DNA sequence as given in any of SEQ ID NOs 1, 3, 5, 7, 9 or 11, or a functional fragment of a nucleic acid as defined in any one of (a) to (m), and,
- (o) nucleic acid sequences encoding a protein as defined in SEQ ID NO 2, 4, 6, 8, 10 or 12 or a nucleic acid as defined in any one of (a) to (n) characterized in that said sequence is a cDNA sequence.
- 3. An isolated nucleic acid according to claim 2 which is DNA, cDNA, genomic DNA or synthetic DNA, or RNA wherein T is replaced by U.
 - 4. A nucleic acid molecule of at least 15 nucleotides in length hybridizing specifically with a nucleic acid of claim 2 or 3.
 - 5. A nucleic acid molecule of at least 15 nucleotides in length specifically amplifying a nucleic acid of claim 2 or 3.
- 6. A vector comprising a nucleic acid sequence according to claim 2 or 3.

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- 7. A vector according to claim 6 which is an expression vector wherein the nucleic acid sequence is operably linked to one or more control sequences allowing the expression of said sequence in prokaryotic and/or eukaryotic host cells.
- 8. A host cell containing a nucleic acid molecule according to claim 2 or 3 or a vector according to claim 6 or 7.
 - 9. The host cell of claim 8, wherein the host cell is a bacterial, insect, fungal, plant or animal cell.
 - 10. An isolated polypeptide encodable by a nucleic acid of claim 2 or 3, or a homologue or a derivative thereof, or an immunologically active and/or functional fragment thereof.

- 11. The polypeptide of claim 10 which has an amino acid sequence as given in SEQ ID NO 2, 4, 6, 8, 10 or 12, or a homologue or a derivative thereof, or an immunologically active and/or functional fragment thereof.
- 12. A method for producing a polypeptide according to claim 10 or 11 comprising culturing a host cell of claim 8 or 9 under conditions allowing the expression of the polypeptide and recovering the produced polypeptide from the culture.
- 13. An antibody specifically recognizing a polypeptide of claim 10 or 11 or a specific epitope thereof.
- 14. A method for the production of transgenic plants, plant cells or plant tissues comprising the introduction of a nucleic acid molecule according to any of claims 2 or 3 in an expressible format or a vector according to claim 6 or 7 in said plant, plant cell or plant tissue.
- 15. A method for the production of altered plants, plant cells or plant tissues comprising the introduction of a polypeptide of claim 10 or 11 directly into a cell, a tissue or an organ of said plant.
- 16. A method for effecting the expression of a polypeptide of claim 10 or 11 comprising the introduction of a nucleic acid molecule of claim 2 or 3 operably linked to one or more control sequences or a vector according to claim 6 or 7 stably into the genome of a plant cell.
- 17. The method of claim 15 or 16 further comprising regenerating a plant from said plant cell.
 - 18. A transgenic plant cell comprising a nucleic acid sequence of claim 2 or 3 which is operably linked to regulatory elements allowing transcription and/or expression of said nucleic acid in plant cells or obtainable by a method of claim 15 or 16.
- 19. The transgenic plant cell of claim 18 wherein said nucleic acid of claim 2 or 3 is stably integrated into the genome of said plant cell.
 - 20. A transgenic plant or plant tissue comprising plant cells of claim 18 or 19.
 - 21. A harvestable part of a plant of claim 20.

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22. The harvestable part of claim 21 which is selected from the group consisting of seeds, leaves, fruits, stem cultures, rhizomes, roots, tubers and bulbs.

- 23. The progeny derived from any of the plants or plant parts of any of claims 20 to 22.
- 24. A method for stimulating root growth comprising expression of a nucleic acid of claim 2 or 3 or comprising expression of another protein that reduces the level of active cytokinins in plants or plant parts.
- 25. A method for enhancing the formation of lateral and adventitious roots comprising expression of a nucleic acid of claim 2 or 3 or comprising expression of another protein that reduces the level of active cytokinins in plants or plant parts.
 - 26. A method for altering root geotropism comprising altering the expression of a nucleic acid of claim 2 or 3 or comprising expression of another protein that reduces the level of active cytokinins in plants or plant parts.
 - 27. A method of any of claims 24 to 26, said method leading to an increase in yield.

- 28. The method of any of claims 24 to 27 wherein said expression of said nucleic acid occurs under the control of a strong constitutive promoter.
- 29. The method of any of claims 24 to 27 wherein said expression of said nucleic acid occurs under the control of a promoter that is preferentially expressed in roots.
 - 30. A method for identifying and obtaining proteins interacting with a polypeptide of claim 10 or 11 comprising a screening assay wherein a polypeptide of claim 10 or 11 is used.
- 31. The method of claim 30 comprising a two-hybrid screening assay wherein a polypeptide of claim 10 or 11 as a bait and a cDNA library as prey are used.
 - 32. A method for modulating the interaction between a polypeptide of claim 10 or 11 and interacting protein partners obtainable by a method according to claim 30 or 31.
 - 33. A method for identifying and obtaining compounds interacting with a polypeptide of claim 10 or 11 comprising the steps of:
- (a) providing a two-hybrid system wherein a polypeptide of claim 10 or 11 and an interacting protein partner obtainable by a method according to claim 30 or 31 are expressed,
 - (b) interacting said compound with the complex formed by the expressed polypeptides as defined in a), and,
- (c) performing measurement of interaction of said compound with said polypeptide or the complex formed by the expressed polypeptides as defined in (a).

- 34. A method for identifying compounds or mixtures of compounds which specifically bind to a polypeptide of claim 10 or 11, comprising:
 - (a) combining a polypeptide of claim 10 or 11 with said compound or mixtures of compounds under conditions suitable to allow complex formation, and,
- (b) detecting complex formation, wherein the presence of a complex identifies a compound or mixture which specifically binds said polypeptide.
 - 35. A method of any of claims 30 to 34 wherein said compound or mixture inhibits the activity of said polypeptide of claim 10 or 11 and can be used for the rational design of chemicals.
- 36. Use of a compound or mixture identified by means of a method of any of claims 30 to 34 as a plant growth regulator or herbicide.

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- 37. A method for production of a plant growth regulator or herbicide composition comprising the steps of the method of any of claims 30 to 34 and formulating the compounds obtained from said steps in a suitable form for the application in agriculture or plant cell or tissue culture.
- 38. A method for the design of or screening for growth-promoting chemicals or herbicides comprising the use of a nucleic acid of claim 2 or 3 or a vector of claim 6 or 7.
- 39. Use of a nucleic acid molecule of claim 2 or 3, the vector of claim 6 or 7, a polypeptide of claim 10 or 11 for increasing yield.
 - 40. Use of a nucleic acid molecule of claim 2 or 3, the vector of claim 6 or 7, a polypeptide of claim 10 or 11 for stimulating root growth.
 - 41. Use of a nucleic acid molecule of claim 2 or 3, the vector of claim 6 or 7, a polypeptide of claim 10 or 11 for enhancing the formation of lateral and adventitious roots.
 - 42. Use of a nucleic acid molecule of claim 2 or 3, the vector of claim 6 or 7, a polypeptide of claim 10 or 11 for altering root geotropism.
 - 43. Diagnostic composition comprising at least a nucleic acid molecule of any of claims 2 to 5, the vector of claim 6 or 7, a polypeptide of claim 10 or 11 or an antibody of claim 13.

- 44. A method for increasing the size of the root meristem comprising expression of a nucleic acid of claim 2 or 3 or comprising expression of another nucleic acid encoding a protein that reduces the level of active cytokinins in plants or plant parts, preferably in roots.
- 45. A method for increasing the root size comprising expression of a nucleic acid of claim 2 or 3 or comprising expression of another nucleic acid encoding a protein that reduces the level of active cytokinins in plants or plant parts, preferably in roots.
 - 46. A method for increasing the size of the shoot meristem comprising downregulation of expression of a nucleic acid of claim 2 or 3, preferably in shoots.
- 47. A method for delaying leaf senescence comprising downregulation of expression of a nucleic acid of claim 2 or 3 in leaves, preferably in senescing leaves.
 - 48. A method for altering leaf senescence comprising expression of a nucleic acid of claim 2 or 3 in senescing leaves.
- 49. A method for increasing leaf thickness comprising expression of a nucleic acid of claim 2 or 3 or comprising expression of another nucleic acid encoding a protein that reduces the level of active cytokinins in plants or plant parts, preferably in leaves.
 - 50. A method for reducing the vessel size comprising expression of a nucleic acid of claim 2 or 3 or comprising expression of another nucleic acid encoding a protein that reduces the level of active cytokinins in vessels.
- 51. A method for increasing the vessel size comprising downregulation of expression of a nucleic acid of claim 2 or 3 in vessels.

- 52. A method for inducing parthenocarpy comprising expression of a nucleic acid of claim 2 or 3 or comprising expression of another nucleic acid encoding a protein that reduces the level of active cytokinins in plants or plant parts, preferably in the placenta, ovules and tissues derived therefrom.
- 53. A method for improving standability of seedlings comprising expression of a nucleic acid of claim 2 or 3 or comprising expression of another protein that reduces the level of active cytokinins in seedlings, preferably in roots.

- 54. A method for increasing branching comprising expression of a nucleic acid of claim 2 or 3 in plants or plant parts, preferably in stems or axillary buds
- 55. A method for improving lodging resistance comprising expression of a nucleic acid of claim 2 or 3 in plants or plant parts, preferably in stems or axillary buds

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ABSTRACT

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The present invention relates to methods for stimulating root growth and/or enhancing the formation of lateral and adventitious roots and/or altering root geotropism comprising expression of a plant cytokinin oxidase or comprising expression of another protein that reduces the level of active cytokinins in plants or plant parts.

The invention also relates to novel plant cytokinin oxidase proteins, nucleic acid sequences encoding cytokinin oxidase proteins as well as to vectors, host cells, transgenic cells and plants comprising said sequences. The invention also relates to the use of said sequences for increasing yield and/or enhancing early vigor and/or modifying root/shoot ratio and/or improving resistance to lodging and/or increasing drought tolerance and/or promoting in vitro propagation of explants and/or modifying cell fate and/or plant development and/or plant morphology and/or plant biochemistry and/or plant physiology. The invention also relates to the use of said sequences in the above mentioned methods.

The invention also relates to methods for identifying and obtaining proteins and compounds interacting with cytokinin oxidase proteins. The invention also relates to the use of said compounds as a plant growth regulator or herbicide.

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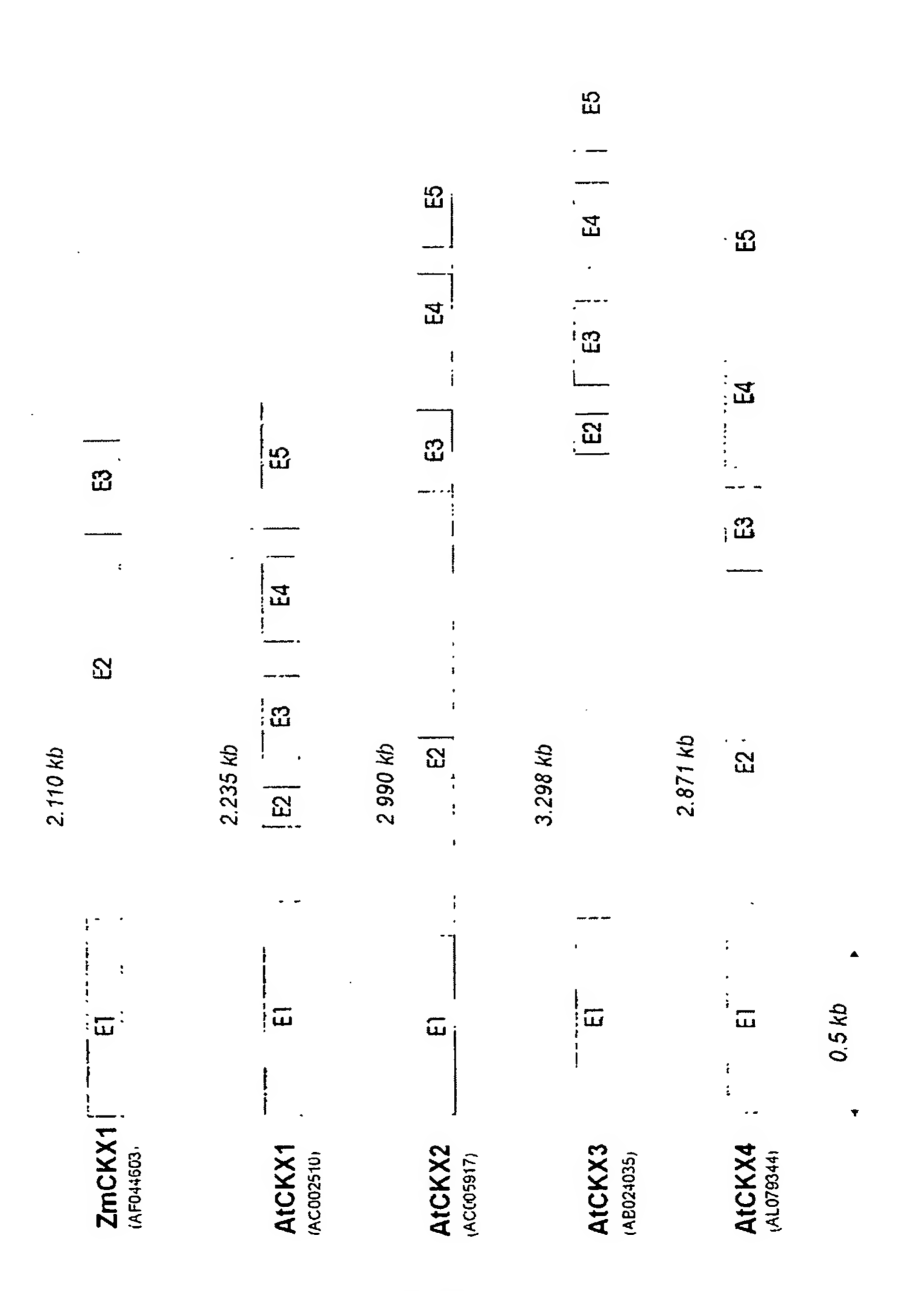


FIGURE 1

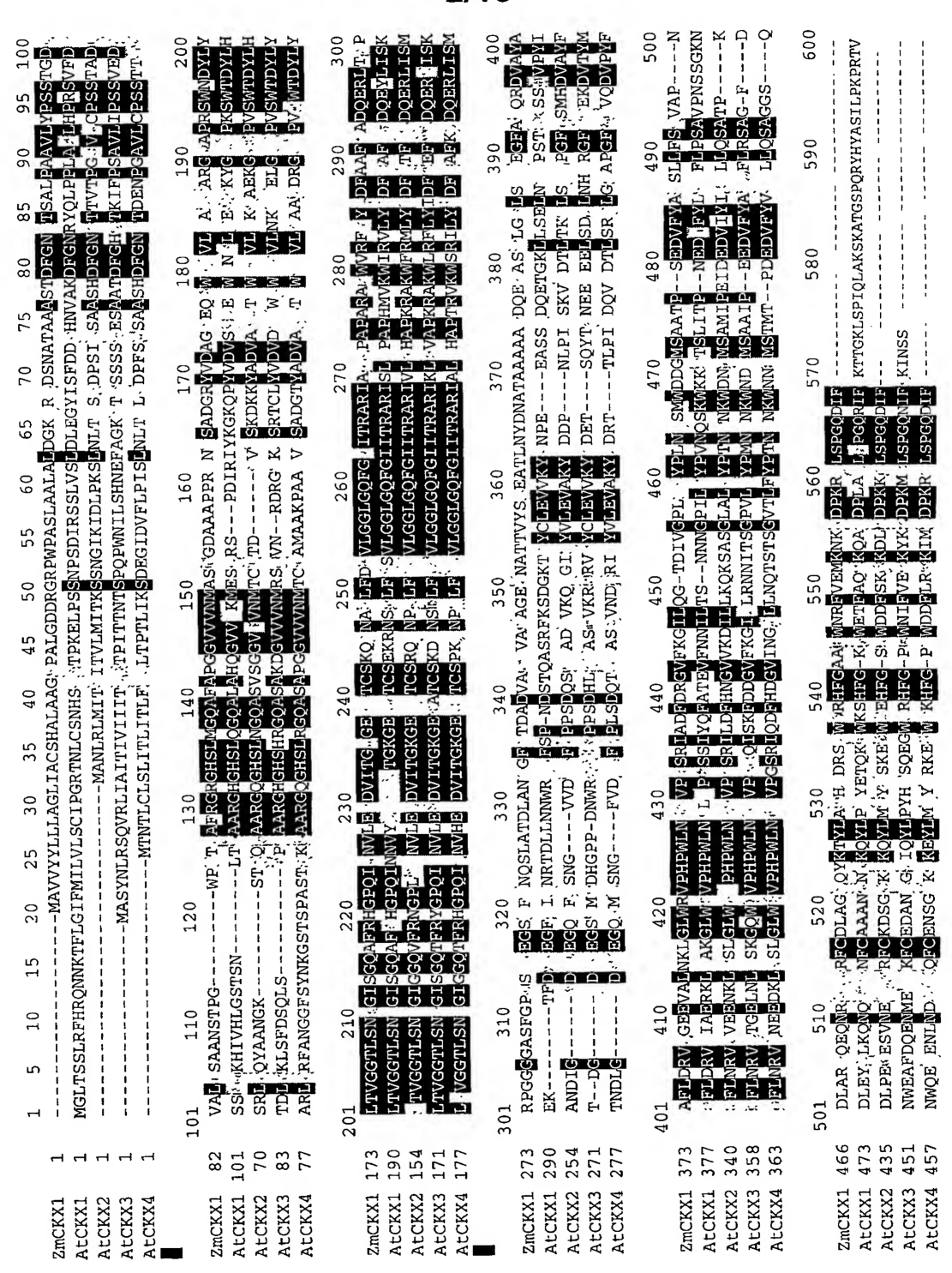


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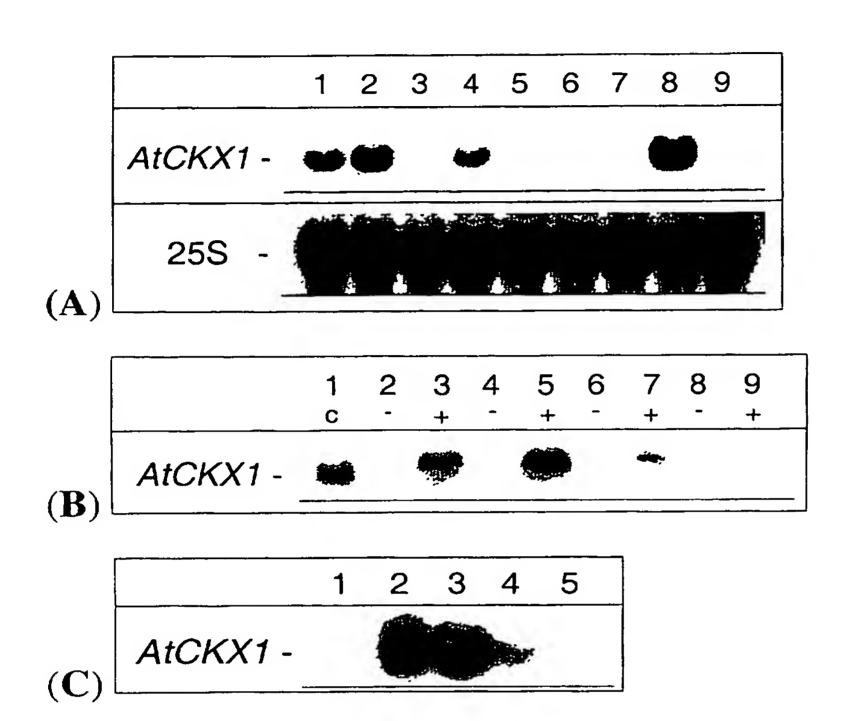


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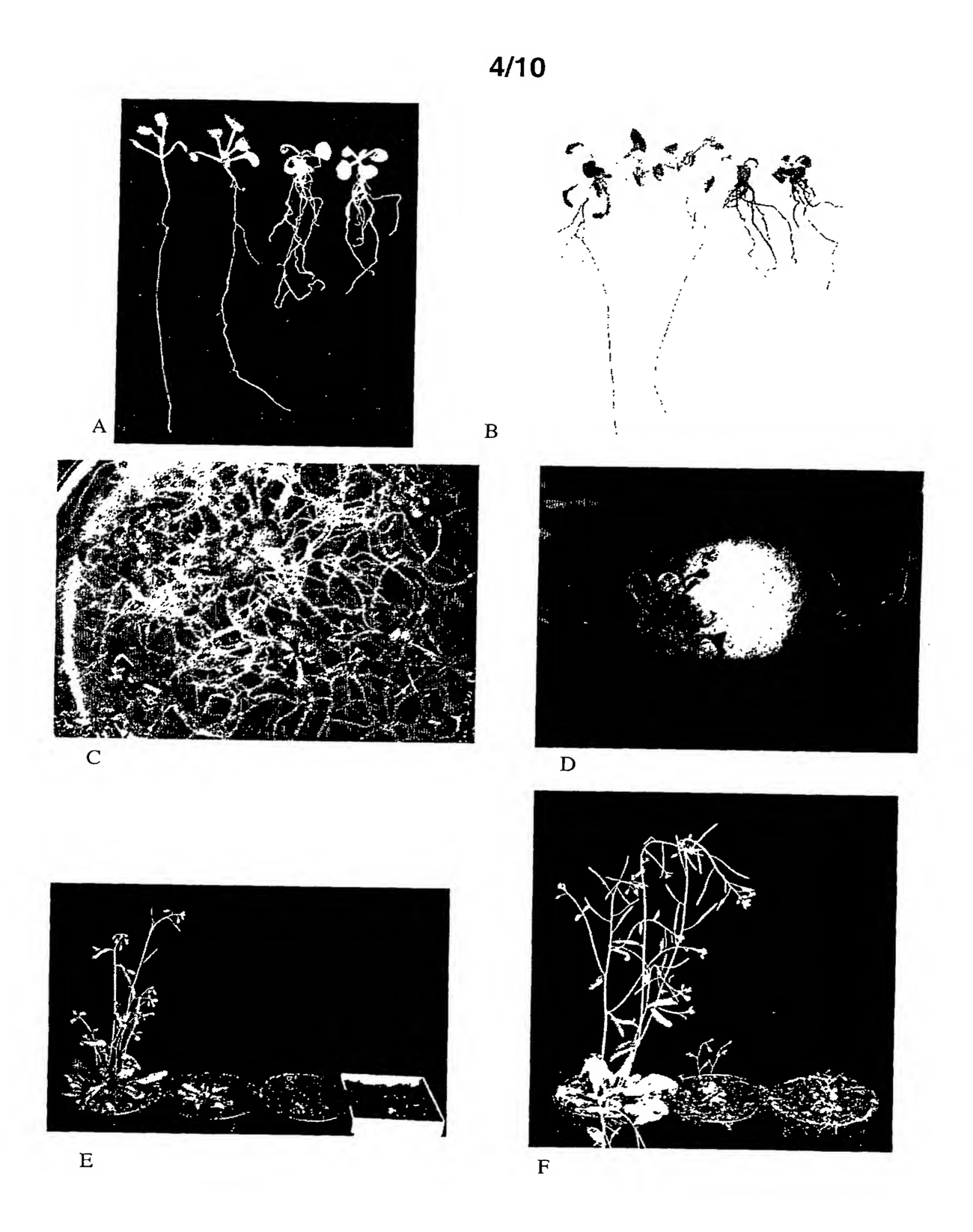


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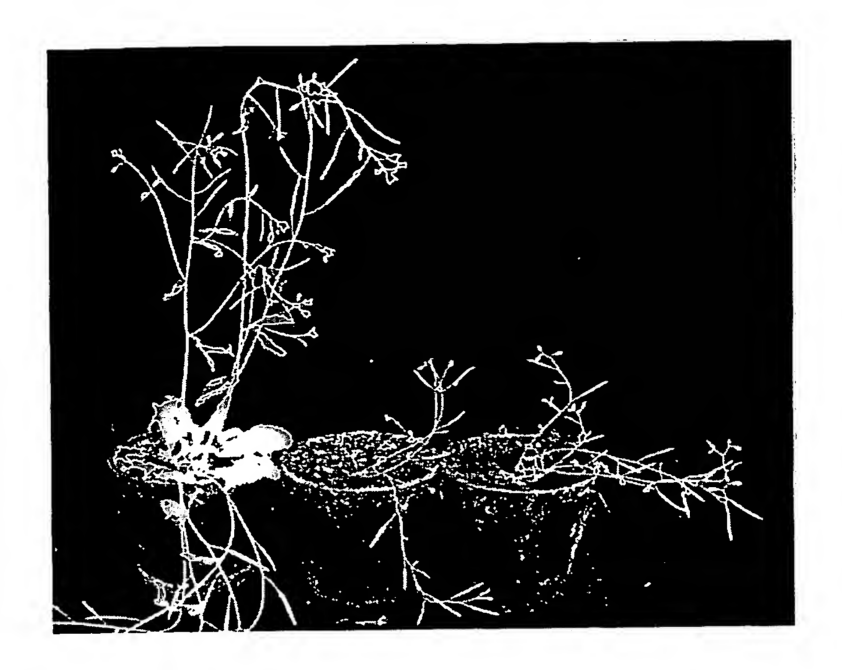


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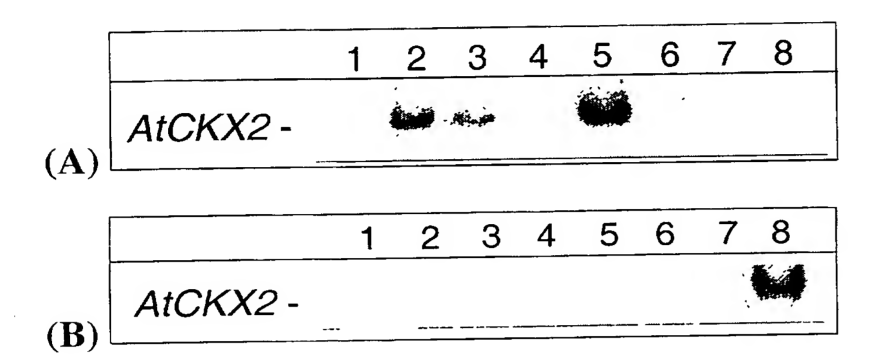


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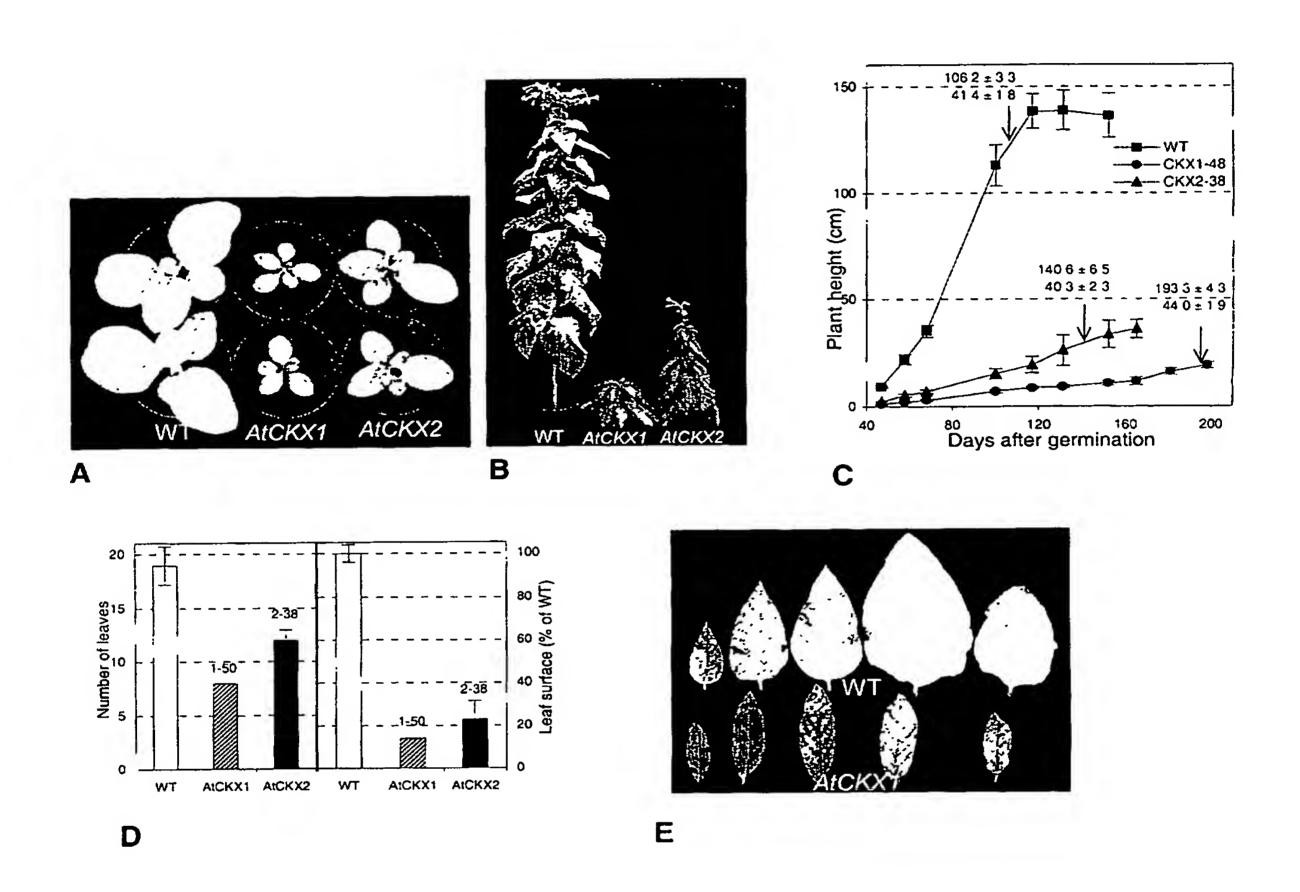
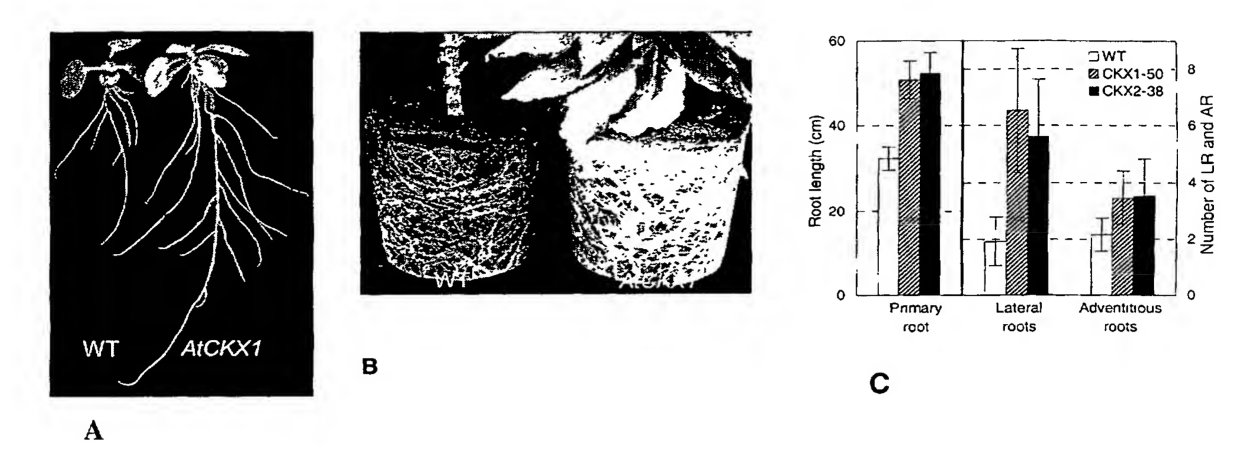


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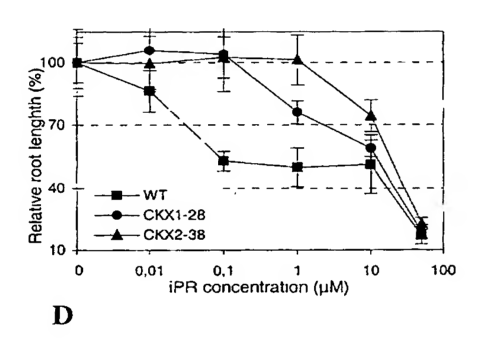


Figure 8



Figure 9

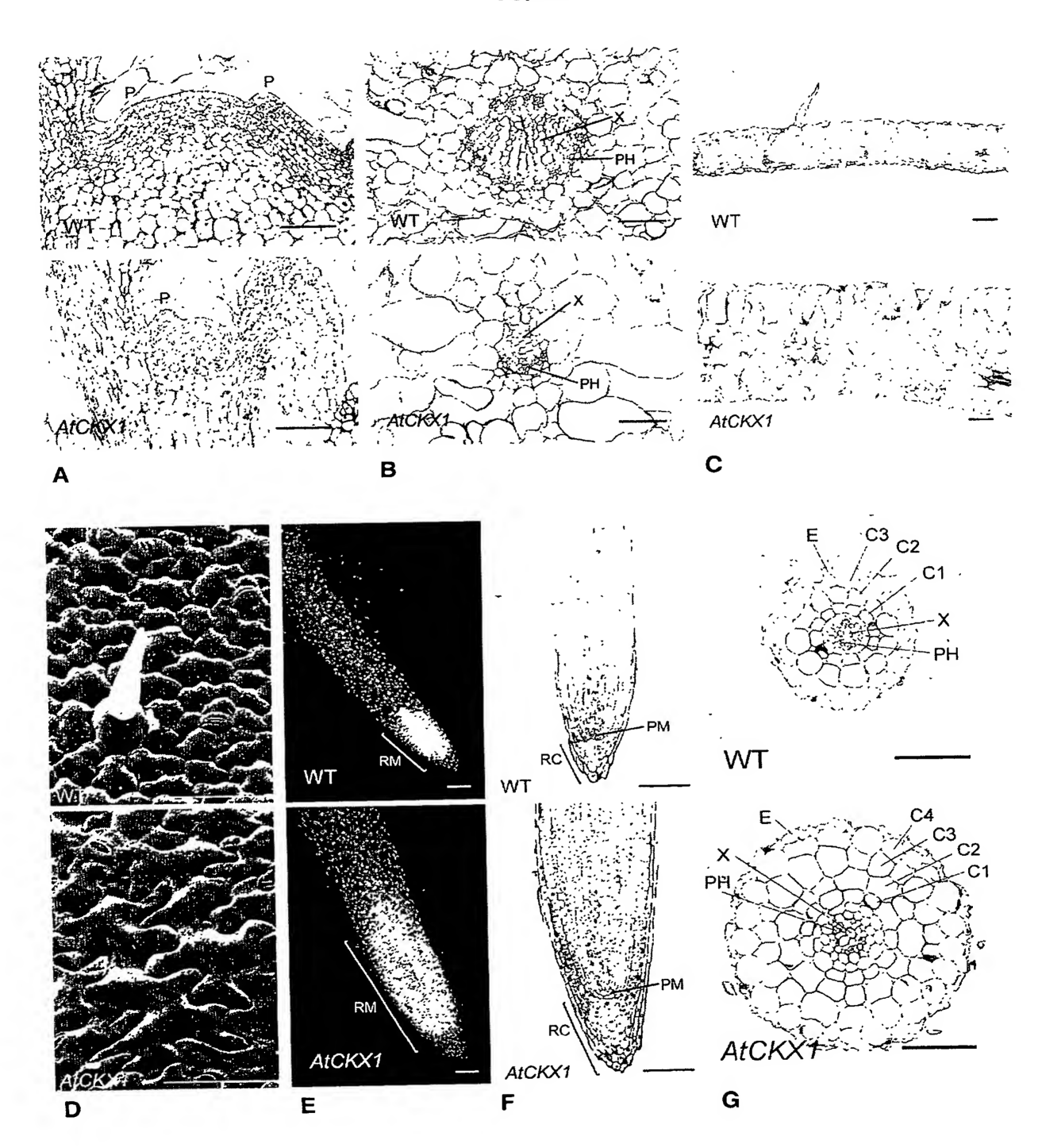


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Val Thr Lys Ile Phe Pro Ser Ala Val Leu Ile Pro Ser Ser Val Glu 65 70 75 80

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Ala Lys Asp Gly Val Val Val Asn Met Arg Ser Met Val Asn Arg Asp

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Ser Ser Gln Glu Gly Trp Val Arg His Phe Gly Pro Arg Trp Asn Ile 485 490 495

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- Gly Gln Ala Ser Ala Pro Gly Gly Val Val Val Asn Met Thr Cys Leu
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- Tyr Ala Asp Val Ala Ala Gly Thr Met Trp Val Asp Val Leu Lys Ala
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- Leu Ser Val Gly Gly Thr Leu Ser Asn Ala Gly Ile Gly Gly Gln Thr
- Phe Arg His Gly Pro Gln Ile Ser Asn Val His Glu Leu Asp Val Ile
- Thr Gly Lys Gly Glu Met Met Thr Cys Ser Pro Lys Leu Asn Pro Glu
- Leu Phe Tyr Gly Val Leu Gly Gly Leu Gly Gln Phe Gly Ile Ile Thr
- Arg Ala Arg Ile Ala Leu Asp His Ala Pro Thr Arg Val Lys Trp Ser
- Arg Ile Leu Tyr Ser Asp Phe Ser Ala Phe Lys Arg Asp Gln Glu Arg
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Val Leu Lys Lys Thr Leu Glu His Gly Leu Ala Pro Lys Ser Trp Thr 145 150 155 160

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Ser Gly Gln Ala Phe His His Gly Pro Gln Ile Ser Asn Val Leu Glu

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Phe Lys Leu Ala Cys Cys Phe Ser Ser Ser Ile Ser Ser Leu Lys Ala 20 25 30

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Lys Asp Phe Gly Asn Arg Tyr Gln Leu Ile Pro Leu Ala Val Leu His
50 55 60

Pro Lys Ser Val Ser Asp Ile Ala Ser Thr Ile Arg His Ile Trp Met 70 75 80

Met Gly Thr His Ser Gln Leu Thr Val Ala Ala Arg Gly Arg Gly His
85 90 95

Ser Leu Gln Gly Gln Ala Gln Thr Arg His Gly Ile Val Ile His Met 100 105 110

Glu Ser Leu His Pro Gln Lys Leu Gln Val Tyr Ser Val Asp Ser Pro 115 120 125

Ala Pro Tyr Val Asp Val Ser Gly Gly Glu Leu Trp Ile Asn Ile Leu 130 135 140

His 145	Glu	Thr	Leu	Lys	Туr 150	Gly	Leu	Ala	Pro	Lys 155	Ser	Trp	Thr	Asp	Tyr 160
Leu	His	Leu	Thr	Val 165	Gly	Gly	Thr	Leu	Ser 170	Asn	Ala	Gly	Ile	Ser 175	Gly
Gln	Ala	Phe	Arg 180	His	Gly	Pro	Gln	Ile 185	Ser	Asn	Val	His	Gln 190	Leu	Glu
Ile	Val	Thr 195	Gly	Lys	Gly	Glu	Ile 200	Leu	Asn	Cys	Thr	Lys 205	Arg	Gln	Asn
Ser	Asp 210	Leu	Phe	Asn	Gly	Val 215	Leu	Gly	Gly	Leu	Gly 220	Gln	Phe	Gly	Ile
Ile 225	Thr	Arg	Ala	Arg	Ile 230	Ala	Leu	Glu	Pro	Ala 235	Pro	Thr	Met	Asp	Gln 240
Glu	Gln	Leu	Ile	Ser 245	Ala	Gln	Gly	His	Lys 250	Phe	Asp	Tyr	Ile	Glu 255	Gly
Phe	Val	Ile	Ile 260	Asn	Arg	Thr	Gly	Leu 265	Leu	Asn	Ser	Trp	Arg 270	Leu	Ser
Phe	Thr	Ala 275	Glu	Glu	Pro	Leu	Glu 280	Ala	Ser	Gln	Phe	Lys 285	Phe	Asp	Gly
Arg	Thr 290	Leu	Tyr	Cys	Leu	Glu 295	Leu	Ala	Lys	Tyr	Leu 300	Lys	Gln	Asp	Asn
Lys 305	Asp	Val	Ile	Asn	Gln 310	Glu	Val	Lys	Glu	Thr 315	Leu	Ser	Glu	Leu	Ser 320
Tyr	Val	Thr	Ser	Thr 325	Leu	Phe	Thr	Thr	Glu 330	Val	Ala	Tyr	Glu	Ala 335	Phe
Leu	Asp	Arg	Val 340	His	Val	Ser	Glu	Val 345	Lys	Leu	Arg	Ser	Lys 350	Gly	Gln
Trp	Glu	Val 355	Pro	His	Pro	Trp	Leu 360	Asn	Leu	Leu	Val	Pro 365	Arg	Ser	Lys
Ile	Asn 370	Glu	Phe	Ala	Arg	Gly 375	Val	Phe	Gly	Asn	Ile 380	Leu	Thr	Asp	Thr
Ser 385	Asn	Gly	Pro	Val	Ile 390	Val	Tyr	Pro	Val	Asn 395	Lys	Ser	Lys	Trp	Asp 400

Asn Gln Thr Ser Ala Val Thr Pro Glu Glu Glu Val Phe Tyr Leu Val 405 410 415 Ala Ile Leu Thr Ser Ala Ser Pro Gly Ser Ala Gly Lys Asp Gly Val 420 425 430 Glu Glu Ile Leu Arg Arg Asn Arg Ile Leu Glu Phe Ser Glu Glu 435 440 445 Ala Gly Ile Gly Leu Lys Gln Tyr Leu Pro His Tyr Thr Thr Arg Glu 450 455 460 Glu Trp Arg Ser His Phe Gly Asp Lys Trp Gly Glu Phe Val Arg Arg 465 470 475 480 Lys Ser Arg Tyr Asp Pro Leu Ala Ile Leu Ala Pro Gly His Arg Ile 485 490 495 Phe Gln Lys Ala Val Ser Tyr Ser 500 <210> 13 <211> 31 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence:oligonucleotide <400> 13 cggtcgacat gggattgacc tcatccttac g 31 <210> 14 <211> 35 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence:oligonucleotide <400> 14

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